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(56) Related Art

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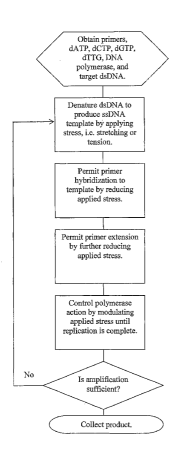
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(54) Title: NANO-PCR: METHODS AND DEVICES FOR NUCLEIC ACID AMPLIFICATION AND DETECTION



(57) Abstract: Methods, devices, and compositions are described that provide for amplification of nucleic acid sequences without reliance upon temperature cycling, thus freeing the methods from conventional benchtop thermal cycling devices. Denaturation of double stranded nucleic acids, primer annealing, and precision control over primer extension by polymerase can be accomplished by applying stress to a nucleic acid. These methods can provide one ore more benefits over conventional PCR methods including: precision control over the PCR process; generally improved fidelity; improved accuracy over problematic sequences such as GC-rich or tandem repeat regions; greater sequence length; increased reaction yield; reduced experimental time; greater efficiency; lower cost; greater portability; and, robustness to various environmental parameters, such as temperature, pH, and ionic strengths.

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For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

NANO-PCR: METHODS AND DEVICES FOR NUCLEIC ACID AMPLIFICATION AND DETECTION

5 Field of the Invention

The invention relates to amplification and detection of nucleic acids. In particular embodiments, the invention provides improved methods, devices, and materials for performing the polymerase chain reaction.

10 Background of the Invention

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The polymerase chain reaction (PCR) has become the conventional technique used to amplify specific DNA or RNA sequences. U.S. Patent No. 4,683,202, issued July 28, 1987 to Mullis and U.S. Patent No. 4,683,195, issued July 28, 1987 to Mullis et al. describe the basic PCR technique. Since the first disclosure of the PCR method, it has had a profound effect on the practice of biotechnology and biomedical science. More than a thousand subsequently-issued U.S. patents reference one or both of these disclosures.

Typically, the amplification of a DNA sequence is performed by first selecting and obtaining two oligonucleotide primers complementary ends of a target DNA sequence. The primers, a polymerase enzyme, a mixture of the four common nucleotide triphosphates, various salts and buffers are mixed with the target DNA which is heated above about 90° C to denature the DNA, separating the target double-stranded DNA into single-stranded DNA templates. Annealing (i.e. sequence-specific hybridization or binding) of the primers to the ends of the DNA templates is promoted by slowly cooling the reaction mixture to less than about 60° C. The temperature is then raised above about 70° C for a period of replication, a process also known as primer extension. The polymerase reads each DNA template strand in the 3' to 5' direction, synthesizing a complementary strand from the ends of the primers in the 5' to 3' direction. This completes one cycle of DNA amplification, which creates starting material for a new cycle. With each complete cycle of denaturation, primer annealing, and primer extension, the process generates

an exponentially increasing (2ⁿ) number of copies of the original, target DNA sequence. To begin a new cycle, the reaction mixture is again heated above 90° C to denature the double-stranded product into single-stranded DNA templates. The primer annealing and extension steps are then repeated.

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This basic PCR amplification scheme, together with various extensions and modifications, enables many different methods for the manipulation and detection of nucleic acids, including for example diagnostic and forensic assays, which require the creation of a threshold amount of DNA from a small initial sample. PCR technology is used, for example, in infectious and genetic disease monitoring, DNA and RNA sequencing, gene expression studies, drug development, and forensic fingerprinting. This has become the standard technology for the detection, identification, and quantification of viral and bacterial pathogens. Several PCRbased diagnostic tests are available for detecting and/or quantifying pathogens, for example, including: HIV-1, which causes AIDS; hepatitis B and C viruses, which can cause liver cancer; human papillomarvirus, which can cause cervical cancer: RSV, which is the leading cause of pneumonia and bronchiolitis in infants: Chlamydia trachomatis and Neisseria gonorrhoeae, which can lead to pelvic inflammatory disease and infertility in women; cytomegalovirus, which can cause life-threatening disease in transplant patients and other immuno-compromised people; and, Mycobacterium tuberculosis, which causes cough and fatigue in its active state and can irreversibly damage infected organs. However, despite addressing needs in numerous areas, current PCR and PCR-based technologies still suffer from several substantial limitations.

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Limitations of Conventional PCR and PCR-Based Technologies

Fidelity: Accuracy on normal sequences limits conventional PCR. For example, Taq, a thermostable polymerase commonly used for DNA amplification, exhibits an error rate of approximately 1 x 10⁻⁴ errors/base pair during PCR. This means that the PCR amplification of a 400 base pair DNA sequence will randomly introduce approximately 40,000 errors among all molecules in the PCR product over 20 cycles.

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Accuracy on difficult target sequences (e.g. GC rich or repeat sequences) is an even more significant limitation of conventional PCR and PCR- based technologies. The error rate for conventional polymerase enzymes such as Taq, depends strongly on the target nucleotide sequence. For example, when the sequence is G+C rich (as seen for example in the 5' regulatory region of the chicken avidin gene), PCR with Taq is oftentimes not a viable process. Likewise, simple repeating sequences, such as trinucleotide repeats (AGC)n or other tandem repeats (A)n, can increase Taq's error rate to 1.5 x 10-2 errors per repeat sequence. See, Shinde et al., Nucleic Acids Research, 31:974. For this reason, several patents have been issued for polymerases that have been genetically engineered to have incrementally higher fidelity (i.e. lower error rates). These include Hi-Fidelity and Phusion Polymerases.

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Length Limitations: The length of the target sequence to be amplified also limits current PCR techniques. Although a few reports have claimed amplification of sequences up to 10 to 20 kilobases, this is highly unusual and quite difficult in routine practice. Moreover, PCR amplification of long target templates is only possible on a limited set of well-behaved DNA sequences. The practical upper-limit for fairly routine and cost-effective amplification of DNA on well-behaved sequences is about 300 to 400 bases in length and is generally reduced for sequences having high G-C content.

Limited Amplification: Current PCR techniques are also limited in the number of amplification cycles that can be carried out in a reaction mixture. Repeated heating and cooling cycles result in progressive enzyme degradation, which limits the factor by which starting material can be amplified. Conventional PCR amplification can rarely be extended beyond 30-35 cycles.

Robustness: Conventional PCR typically requires significant volumes of reagents, bulky equipment (e.g., thermal cyclers), substantial human labor (e.g., tedious optimizations), and minimum amounts of starting material, each of which contributes to making conventional PCR a costly and time-consuming process. Current PCR techniques typically take from several hours for normal sequences to

several days to weeks for difficult sequences or long template. Conventional PCR

- 4 -

requires a significant amount of time to cycle and equilibrate the temperature of the reaction mix. Moreover, time-consuming optimizations can be required in order to reliably amplify targets that are less than ideal.

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Tightly controlled conditions (*e.g.*, temperature, pH, and buffer ingredients) are required for performance of conventional PCR techniques. Additionally, various contaminants can interfere with PCR amplification by directly inhibiting or interfering with polymerase enzymes used to copy the target DNA or RNA. This further limits the quality of starting material that can be used for amplification and places additional requirements on the level of purity that must be obtained by DNA or RNA extraction techniques before the amplification steps can be reliably performed. The performance environment of conventional PCR is generally limited to laboratories, and is rarely practicable in remote locations, doctor's offices, at the patient's bedside, or out in the field.

Sensitivity and Specificity of Diagnostics: The sensitivity of PCR-based diagnostic and forensic kits and assays depends on the overall yield, accuracy, robustness, and target length achievable in a PCR reaction. The above-mentioned limitations in performance parameters of current PCR set limits on the minimum amount of starting DNA or RNA necessary in order to reliably carry out PCR amplification. This, in turn, limits the sensitivity of any pathogen detection system, diagnostic, or forensic kits or assays that rely upon conventional PCR or PCR-based technologies. The specificity of a PCR-based diagnostic, forensic, or pathogen detection system depends critically on the accuracy with which DNA can be amplified and read as well as the length of the target DNA or RNA that can be reliably amplified and identified.

For these and other reasons, current generation PCR-based technologies and detection systems are generally limited with respect to overall speed, efficiency, cost-effectiveness, and scope of use. Incremental improvements to conventional PCR methods and devices have been proposed with respect to some of the isolated performance parameters described above. For example, Tso et al. discloses a PCR microreactor for amplifying DNA using microquantities of sample fluid in U.S. Patent No. 6,613,560, issued September 2, 2003. Alternatives to high temperature

-5-(followed by page 5a)

DNA denaturation have also been proposed. For example, Purvis disclosed a method of electrochemical denaturation of double-stranded nucleic acid in U.S. Patent No. 6,291,185, issued September 18, 2001. Stanley discloses another method of electrochemical denaturation of nucleic acids in U.S. Patent No. 6,197,508, issued March 6, 2001. Dattagupta et al. have disclosed a method of using primers to displace the DNA strand from the template in U.S. Patent No. 6,214,587, issued April 10, 2001. Mullis, *supra*, suggested the use of helicase enzymes for separating DNA strands.

In view of the limitations of conventional PCR, and despite the proposal of various incremental improvements, there remains a need in the art for improved methods, devices, and compositions for the amplification, manipulation, sequencing, and detection of nucleic acids. It is therefore an object of the present invention to go someway towards meeting this need; or to at least provide the public with a useful choice.

In this specification where reference has been made to patent specifications, other external documents, or other sources of information, this is generally for the purpose of providing a context for discussing the features of the invention. Unless specifically stated otherwise, reference to such external documents is not to be construed as an admission that such documents, or such sources of information, in any jurisdiction, are prior art, or form part of the common general knowledge in the art.

Summary of the Invention

The methods and apparatuses described herein provide a breakthrough technology to perform PCR. The technology described herein also permits PCR to be performed without reliance upon thermal cycling. The technology may be applied at a wide range of ambient temperatures or using controlled temperature. It is possible to exercise precise control when desired during replication and amplification, thereby enabling substantial improvements in a number of performance parameters. Dubbed "Nano-PCRTM," this technology introduces a new paradigm in PCR-based detection and amplification of nucleic acids.

--5a-(followed by page 5b)

In one aspect the present invention provides a method for amplifying a nucleic acid, comprising:

- (a) contacting one or more template strands of single-stranded nucleic acid with one or more oligonucleotide primers complementary to a portion of the one or more template strands:
- (b) annealing at least one primer of the one or more primers to the portion of the one or more template strands to which the primer is complementary;
- (c) contacting the one or more template strands with a nucleic acid polymerase and at least four different nucleoside triphosphates;
- (d) extending the at least one annealed primer by the nucleic acid polymerase thereby forming one or more extension products bound to the one or more template strands;
- (e) separating the one or more extension products from the one or more template strands; and
- (f) repeating steps (a), (b), (c), (d) and (e) to amplify the nucleic acid,

wherein at least one of the one or more extension products in step (e) is used as template strands in a subsequent cycle of steps (a)-(e), wherein, for the last cycle, step (c) is optional, and wherein at least one of steps (b), or (d) comprises applying tension to the one or more template strands.

In another aspect the present invention provides a method for amplifying a nucleic acid, comprising:

- contacting one or more template strands of single-stranded. (a) nucleic acid with one or more oligonucleotide primers complementary to a portion of the one or more template strands, wherein at least one primer of the one or more primers comprises one or more complexing groups capable of binding one or more molecules bound to a substrate;
- (b) annealing the at least one primer with the one or more complexing groups to the portion of the one or more template

-5b-(followed by page 5c)

strands to which the primer is complementary, thereby binding the one or more template strands to the substrate;

- (c) contacting the one or more template strands with a nucleic acid polymerase and at least four different nucleoside triphosphates;
- extending the one or more primers by the nucleic acid polymerase thereby forming one or more extension products bound to the one or more template strands;
- separating the one or more extension products from the one or more template strands; and
- (f) repeating steps (a), (b), (c), (d) and (e) to amplify the nucleic acid, wherein, in the last cycle, step (e) is optional, wherein at least one of the one or more extension products in step (e) is used as template strands in a subsequent cycle of steps (a)-(e), and

wherein at least one of steps (b), (d), and (e) comprises applying tension to the one or more template strands.

In another aspect the present invention provides a method of detecting a pathogen comprising the method of the invention and further comprising probing the amplified nucleic acid for the presence of a nucleotide sequence specifically associated with the pathogen.

In another aspect the present invention provides a method for amplifying a nucleic acid, comprising:

- (a) denaturing a double-stranded nucleic acid molecule by applying tension to one or both strands of the double-stranded nucleic acid molecule sufficient to cause separation of the double-stranded nucleic acid molecule into two single-stranded nucleic acid molecules, wherein the one or both single-stranded nucleic acid molecules to which the tension is applied becomes a template strand;
- (b) contacting the one or both template strands with one or more oligonucleotide primers complementary to a portion of the

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-5c-(followed by page 5d)

template strands, a nucleic acid polymerase and at least four different nucleoside triphosphates;

- (c) reducing the tension applied to the one or both template strands in step a) to permit the one or more primers to anneal to the corresponding portions of the template strands to which the one or more primers are complementary;
- (d) controlling the tension applied to the one or both template strands so as to permit extension of the one or more primers by the nucleic acid polymerase, thereby forming one or more double-stranded nucleic acid molecules in which one or more extension products are bound to the one or more template strands; and
- (e) repeating steps (a), (b), (c), and (d) to amplify the nucleic acid, wherein at least one of the one or more double-stranded nucleic acid molecules formed in step (d) in a cycle of steps (a)-(d) is used as the double-stranded nucleic acid molecule in step (a) in a subsequent cycle of steps (a)-(d), so that at least one extension product is formed in step (d) is used as a template strand in a subsequent cycle of steps (a)-(d).

In another aspect the present invention provides a method of performing nucleic acid amplification, comprising:

- (a) denaturing a double-stranded nucleic acid to produce singlestranded template strands;
- (b) annealing at least one primer to the template strands;
- (c) extending at least one primer by a template-strand-driven polymerase to produce extension products, thereby forming double-stranded nucleic acid molecules; and
- (d) repeating steps (a)-(c) to amplify the double-stranded nucleic acid,

wherein at least one of the extension products in step (c) is used as template strand in a subsequent cycle of steps (a)-(c), and wherein step (c) comprises applying adjustable tension to the nucleic acid, said adjustable tension modulating proofreading exonuclease activity of the nucleic acid polymerase.

-5d-(followed by page 5e)

In another aspect the present invention provides a method for controlling the stringency of annealing of a primer to a nucleic acid strand during nucleic acid amplification, comprising:

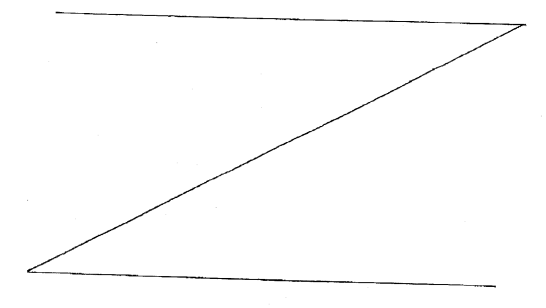
annealing the primer to the nucleic acid strand, wherein tension is adjustably applied to said nucleic acid strand or to said primer to control the stringency of annealing of said primer to said nucleic acid strand.

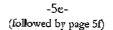
In another aspect the present invention provides a method for lowering the error rates of extending a primer by a nucleic acid polymerase during nucleic acid amplification, comprising:

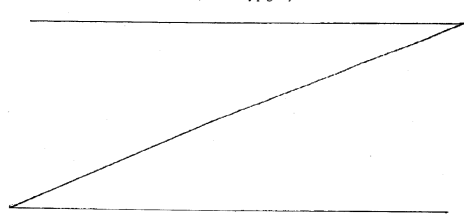
extending the primer annealed to the nucleic acid template by a nucleic acid polymerase to produce double-stranded nucleic acid molecules, wherein tension is adjustably applied to said nucleic acid template or to said primer to lower the error rates of extending said primer by the nucleic acid polymerase.

In another aspect the present invention provides a method for lowering the error rates of extending a primer on a GC-rich template by a nucleic acid polymerase during nucleic acid amplification, comprising:

extending the primer annealed to the GC-rich nucleic acid template by a nucleic acid polymerase to produce double-stranded nucleic acid molecules, wherein tension is adjustably applied to said GC-rich nucleic acid template or to said primer to lower the error rates of extending said primer on said GC-rich template.







In another aspect the present invention provides a method of improving fidelity of nucleic acid sequencing, comprising:

- (a) providing four samples, each sample comprising:
 - a single-stranded or a double-stranded DNA target sequence;
 - a DNA polymerase;
 - a primer complementary to the target sequence;
 - at least four different nucleoside triphosphates; and
 - a respective dideoxy nucleoside triphosphate selected from among ddATP, ddCTP, ddGTP, and ddTTP;
- (b) denaturing the double-stranded DNA target sequence into single-stranded DNA target sequence;
- (c) hybridizing the primer to its complementary single-stranded DNA target sequence;
- (d) extending the primers to form double-stranded DNA extension products while adjustably applying tension to the template strand to induce proofreading exonuclease activity, thereby improving fidelity or primer extension;
- (e) repeating steps (b) through (d) to form further extension products, wherein at least one of the extension products formed in step (d) is used as a template strand in a subsequent cycle of steps (b)-(d); and
- (f) determining the sequence of the DNA target sequence, wherein more than one of the steps (b) through (d) comprises applying tension to one or more strands of DNA target sequence.

-5f-(followed by page 5g)

In another aspect the present invention provides a product produced by the method of the invention.

In another aspect the present invention provides a method of detecting presence of a pathogen in a biological sample, comprising:

- (a) performing a nucleic acid amplification:
 - (i) denaturing a double-stranded nucleic acid to produce singlestranded template strands;
 - (ii) annealing at least one primer to the template strands;
 - (iii) extending the at least one primer by a template-stranddriven polymerase to produce extension products, thereby forming double-stranded nucleic acid molecules; and
- (iv) repeating steps (i)-(iii) to amplify the double-stranded nucleic acid,

wherein at least one of steps (i)-(iii) comprises applying tension to the nucleic acid, and

(b) probing the amplified nucleic acid for the presence of a nucleotide sequence specifically associated with the pathogen, wherein pathogen or the nucleotide sequence are detected at concentrations lower than 1000 organisms or polynucleotides/ml.

In another aspect the present invention provides a method of detecting presence of at least one specific nucleic acid sequence in a sample, comprising:

- (a) performing a nucleic acid amplification:
 - (i) denaturing a double-stranded nucleic acid to produce single-stranded template strands;
 - (ii) annealing at least one primer to the template strands;
 - (iii) extending the at least one primer by a template-stranddriven polymerase to produce extension products, thereby forming double-stranded nucleic acid molecules; and

-5g-(followed by page 5h)

(iv) repeating steps (i)-(iii) to amplify the double-stranded nucleic acid, thereby producing amplified nucleic acid,

wherein at least one of steps (i)-(iii) comprises applying tension to the nucleic acid, and

- (b) contacting the amplified nucleic acid with at least one probe; and
- (c) detecting whether the specific nucleic sequence is in the sample by observing the presence or absence of the probe bound to the amplified nucleic acid.

In another aspect the present invention provides a method of distinguishing between at least two specific nucleic acid sequences in a sample, comprising:

- (a) performing a nucleic acid amplification:
 - (i) denaturing a double-stranded nucleic acid to produce singlestranded template strands;
 - (ii) annealing at least one primer to the template strands;
 - (iii) extending the at least one primer by a template-stranddriven polymerase to produce extension products, thereby forming double-stranded nucleic acid molecules; and
 - (iv) repeating steps (i)-(iii) to amplify the double-stranded nucleic acid, thereby producing amplified nucleic acid.

wherein at least one of steps (i)-(iii) comprises applying tension to the nucleic acid, and

- (b) contacting the amplified nucleic acid with a plurality of probes; and
- (c) detecting which of a plurality of probes is bound to the amplified nucleic acid, thereby distinguishing between at least two specific nucleic acid sequences.

In another aspect the present invention provides a method of amplification or detection of sequences encoded on RNA, comprising:

(a) isolating RNA from a sample;

-5h-(followed by page 5i)

- (b) performing reverse transcription thereby obtaining a corresponding cDNA template;
- (c) amplifying the corresponding cDNA template by a method, including:
 - (i) optionally denaturing a double-stranded cDNA template to produce at least one single-stranded cDNA template strand;
 - (ii) annealing at least one primer to the at least one cDNA template strand;
 - (iii) extending the at least one primer by a template-stranddriven polymerase to produce extension products, thereby forming double-stranded nucleic acid molecules; and
 - (iv) repeating steps (i)-(iii) to amplify the double-stranded nucleic acid, thereby amplifying or detecting sequences encoded on RNA.

wherein at least one of steps (i)-(iii) comprises applying tension to the nucleic acid.

In another aspect the present invention provides a method for controlling denaturing of a double stranded nucleic acid during nucleic acid amplification, comprising adjustably applying hydrodynamic tension to said nucleic acid strand to control the denaturing of said double stranded nucleic acid.

In another aspect the present invention provides a product of nucleic acids produced by the method of the invention, wherein

- a. the average nucleic acid length can be up to 20,000 base pairs; or
- b. the nucleic acid product contains less than 10⁻⁷ errors /base for normal sequences; or
- c. the nucleic acid product contains less than 10⁻³ errors/base for difficult sequences; or
- d. the nucleic acid product is produced by more than 100 cycles of amplification.

-5i-(followed by page 5j)

In another aspect the present invention provides a method for amplifying a nucleic acid, comprising:

- (a) contacting one or more template strands of single-stranded nucleic acid with one or more oligonucleotide primers complementary to a portion of the one or more template strands;
- (b) annealing at least one primer of the one or more primers to the portion of the one or more template strands to which the primer is complementary;
- (c) contacting the one or more template strands with a nucleic acid polymerase and at least four different nucleoside triphosphates;
- (d) extending the at least one annealed primer by the nucleic acid polymerase, thereby forming one or more extension products bound to the one or more template strands;
- (e) separating the one or more extension products from the one or more template strands; and
- (f) repeating steps (a), (b), (c), (d) and (e) to amplify the nucleic acid,

wherein at least one of the one or more extension products in step (e) is used as template strands in a subsequent cycle of steps (a)-(e),

wherein, for the last cycle, step (e) is optional, and wherein tension is applied during each cycle of steps (a)-(e), and, optionally, wherein each step (a) through (e) is isothermal.

In another aspect the present invention provides a method for improving fidelity of a nucleic acid amplification, comprising:

- (a) contacting one or more template strands of single-stranded nucleic acid with one or more oligonucleotide primers complementary to a portion of the one or more template strands;
- (b) annealing at least one primer of the one or more primers to the portion of the one or more template strands to which the primer is complementary;

-5j-(followed by page 6)

- (c) contacting the one or more template strands with a nucleic acid
 polymerase and at least four different nucleoside triphosphates;
- (d) extending the at least one annealed primer by the nucleic acid polymerase, thereby forming one or more extension products bound to the one or more template strands;
- separating the one or more extension products from the one or more template strands; and
- repeating steps (a) through (e) to amplify the nucleic acid,
 wherein at least one of the one or more extension products in
 step (e) is used as template strands in a subsequent cycle of steps
 (a)-(e) and wherein, for the last cycle, step (e) is optional; and
 wherein

either

tension is applied during each cycle of steps (a)-(e);

or

wherein tension induces proofreading exonuclease activity; or

wherein tension is applied to control at least one of the step (a) through (e),

thereby improving fidelity of amplification.

Brief Description of the Drawings

FIGS. 1A and 1B illustrate exemplary flow charts of PCR methods that do not rely on temperature cycling.

FIGS. 2A-2C illustrate exemplary methods of and arrangements of elements of a reaction chamber for applying tension to a DNA strand anchored between opposed surfaces.

FIGS. 3A and 3B illustrate methods of and arrangements of elements of a reaction chamber for applying tension to a DNA strand using optical or magnetic traps.

FIG. 4 is an illustration of an exemplary method of and arrangement of elements of a reaction chamber for applying tension to a DNA strand bound to a polymerase fixed to a substrate in a fluid flow.

FIGS. 5A and 5B illustrate exemplary methods of and arrangements of elements of a reaction chamber for applying tension to a DNA strand anchored at one end in a fluid flow.

FIGS. 6A-6C illustrate exemplary methods of and arrangements of elements of a reaction chamber for applying tension to a DNA strand suspended in a fluid velocity gradient.

FIGS. 7A and 7B illustrate schematics of exemplary devices for performing a PCR method that does not rely on temperature cycling or thermal denaturation.

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Detailed Description of Embodiments of the Invention

Definitions

The terms "nucleic acid," "polynucleotide," and "oligonucleotide" are used herein to include a polymeric form of nucleotides of any length, including, but not limited to, ribonucleotides or deoxyribonucleotides. There is no intended distinction in length between these terms. Further, these terms refer only to the primary structure of the molecule. Thus, in certain embodiments these terms can include triple-, double- and single-stranded DNA, as well as triple-, double- and single-stranded RNA. They also include modifications, such as by methylation and/or by capping, and unmodified forms of the polynucleotide. More particularly, the terms "nucleic acid," "polynucleotide," and "oligonucleotide," include polydeoxyribonucleotides (containing 2-deoxy-D-ribose), polyribonucleotides (containing D-ribose), any other type of polynucleotide which is an N- or C-glycoside of a purine or pyrimidine base, and other polymers containing nonnucleotidic backbones, for example, polyamide (e.g., peptide nucleic acids (PNAs)) and polymorpholino (commercially available from Anti-Virals, Inc.

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Corvallis, Oreg., as Neugene) polymers, and other synthetic sequence-specific nucleic acid polymers providing that the polymers contain nucleobases in a configuration which allows for base pairing and base stacking, such as is found in DNA and RNA.

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As used herein, "primer" refers to a single-stranded polynucleotide capable of acting as a point of initiation of template-directed DNA synthesis under appropriate conditions (i.e., in the presence of four different nucleoside triphosphates and an agent for polymerization, such as, DNA or RNA polymerase or reverse transcriptase) in an appropriate buffer and at a suitable temperature. The appropriate length of a primer depends on the intended use of the primer but typically is at least 7 nucleotides long and, more typically range from 10 to 30 nucleotides in length. Other primers can be somewhat longer such as 30 to 50 nucleotides long. PCR primers are typically about 20-30 base pairs long and are chosen to be complementary to one strand upstream (i.e., 5' to 3') of the target sequence and the opposite strand downstream (i.e., 3' to 5') of the sequence. The 5' ends of the primers define the ends of the amplified PCR product. Primers may contain approximately the same GC content as AT content and no long stretches of any one base. Furthermore, the primers should not contain structures that are substantially complementary to one another. This insures that "primer dimer" formation or other secondary structure does not occur. Short primer molecules generally require cooler temperatures to form sufficiently stable hybrid complexes with the template. A primer need not reflect the exact sequence of the template but must be sufficiently complementary to hybridize with a template. The term "primer site" or "primer binding site" refers to the segment of the target DNA to which a primer hybridizes. The term "primer pair" means a set of primers including a 5' "upstream primer" that hybridizes with the complement of the 5' end of the DNA sequence to be amplified and a 3' "downstream primer" that hybridizes with the 3' end of the sequence to be amplified.

As used herein, the term "complementary" means that one nucleic acid is identical to, or hybridizes selectively to, another nucleic acid molecule. Selectivity of hybridization exists when hybridization occurs that is more selective than total

lack of specificity. Typically, selective hybridization will occur when there is at least about 55% identity over a stretch of at least 14-25 nucleotides, alternatively at least 65%, at least 75%, or at least 90%. In one alternative embodiment, one nucleic acid hybridizes specifically to the other nucleic acid. See M. Kanehisa, Nucleic Acids Res. 12:203 (1984).

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A primer that is "perfectly complementary" has a sequence fully complementary across the entire length of the primer and has no mismatches. The primer is typically perfectly complementary to a portion (subsequence) of a target sequence. A "mismatch" refers to a site at which the nucleotide in the primer and the nucleotide in the target nucleic acid with which it is aligned are not complementary. The term "substantially complementary" when used in reference to a primer means that a primer is not perfectly complementary to its target sequence; instead, the primer is only sufficiently complementary to hybridize selectively to its respective strand at the desired primer-binding site.

As used herein, a "probe" is a nucleic acid capable of binding to a target nucleic acid of complementary sequence through one or more types of chemical bonds, usually through complementary base pairing, usually through hydrogen bond formation, thus forming a duplex structure. A probe binds or hybridizes to a "probe binding site." A probe can be labeled with a detectable label to permit facile detection of the probe, particularly once the probe has hybridized to its complementary target. A label attached to the probe can include any of a variety of different labels known in the art that can be detected by chemical or physical means, for example. Labels that can be attached to probes include, but are not limited to, radioisotopes, fluorophores, chromophores, gold particles, quantum dots, mass labels, electron dense particles, magnetic particles, spin labels, molecules that emit chemiluminescence, electrochemically active molecules, enzymes, cofactors, and enzyme substrates. Probes can vary significantly in size. Some probes are relatively short. Generally, probes are at least 7 to 15 nucleotides in length. Other probes are at least 20, 30 or 40 nucleotides long. Still other probes are somewhat longer, being at least 50, 60, 70, 80, 90 nucleotides long. Yet other probes are longer still, and are

at least 100, 150, 200 or more nucleotides long. Probes can be of any specific length that falls within the foregoing ranges as well.

A "thermophilic DNA polymerase" is a thermostable DNA polymerase enzyme having an optimum temperature at which it functions, which is higher than 40°C. Oftentimes, the optimum temperature for the function of a thermophilic DNA polymerase ranges from about 50°C to 80°C, or 60°C to 80°C. These heat stable enzymes were introduced to provide more robustness to the repeated cycles of heating and cooling the enzyme during conventional PCR.

A "difficult sequence" refers to sequences on which a polymerase enzyme has a tendency to slip, make mistakes or stop working. Examples of difficult sequences include sequences of several residues (e.g. segments of 6, 9, 12, 15, or 30 base pairs or longer) having greater than about 50% G and C base pairs that are called GC-rich sequences, sequences containing tandem repeat segments, polyrepeat sequences such as poly—A sequences, trinucleotide repeat regions as found in sequences associated certain diseases like Huntington's, and other such problematic sequences.

The term "comprising" as used in this specification means "consisting at least in part of". When interpreting each statement in this specification that includes the term "comprising", features other than that or those prefaced by the term may also be present. Related terms such as "comprise" and "comprises" are to be interpreted in the same manner.

Overview of the Conventional Polymerase Chain Reaction (PCR)

To perform the standard thermal cycling polymerase chain reaction using a thermophilic (i.e. heat stable) DNA polymerase, one typically executes the following steps: 1) prepare a cocktail containing a PCR buffer, a dNTP mixture, a primer pair, a DNA polymerase, and doubly-deionized water in a tube; 2) add the DNA to be amplified to the tube 3) place the tube in a temperature block of a thermal cycler (e.g. Perkin-ElmerTM 9600 or 9700 PCR Thermal Cycler) 4) Program the thermal cycler with specific reaction conditions (e.g. a period for thermal denaturation of double-stranded DNA by heating to above about 90°C for about 1 to 2 minutes, a period of annealing by slowly cooling to about 50 to 65° C for 2 min, and a period for polymerization, also called primer extension, by heating to about 70 to 75° C for a few minutes) that are to be repeated for about 25 to 35 cycles. Executing the method produces about a 2^n fold amplification of the starting material, where n is the total number of cycles of amplification that are carried out.

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While some limitations of conventional PCR stem from how the conventional technique is typically carried out, several limitations in the performance parameters stem directly or indirectly from the reliance on thermal cycling. Overall reaction yield, amplification efficiency, sensitivity, robustness, and portability are each, for instance, restricted by thermal cycling. The Nano-PCRTM methods overcome not only limitations due to thermal cycling but also several that are inherent to typical implementation of conventional PCR.

Nano-PCRTM

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Nano-PCRTM methods and apparatuses can dramatically extend the detection and amplification capabilities of the polymerase chain reaction by breaking through several limitations imposed by conventional approaches. Table 1 compares typical performance parameters of current PCR with Nano-PCRTM.

Table 1: Performance parameters of current PCR vs. Nano-PCRTM

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Performance Parameters	Current Generation PCR	Nano-PCRTM	
		Methods and Devices	
Accuracy-normal sequences	Typical error rate of about 1 x	Error rates of less than about	
	10 ⁻⁴ errors/base pair.	1.0 x 10 ⁻⁷ errors/base pair or	
		better can be achieved.	
Accuracy when replicating	Typical error rate of about 1.5	Error rates of less than about	
problematic sequences (GC	x 10 ⁻² errors/repeat sequence.	1.0 x 10 ⁻³ errors/repeat	
rich or repeating sequences)		sequence can be achieved with	
		precision control of the	
		replication process.	
Length of amplified sequence	Typically limited to about	Can amplify extended	
	300-400 base pairs.	sequences of up to 20,000 base	
		pairs or more.	
Overall reaction yield	Reagent and polymerase	Reagents can survive more	
	degradation generally limits	than 100 cycles of DNA	
	amplification to about 30-35	amplification.	
	cycles.		

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Cost	Thermal cycler generally	Small, relatively inexpensive
	costs more than about \$4,000.	devices can be manufactured
		to perform Nano-PCR TM .
Overall PCR process time	Can require several hours to	Can be performed in less
	days.	than 1 hour.
Portability	Performed in bench top	Can be performed in a
	devices that are restricted to	portable hand-held device.
	laboratory settings.	
Robustness	Requires tightly controlled	Allows for temperature and
	operational conditions i.e.	pH variations. Can work with
	temperature, pH. Need	a broader range of starting
	Highly Pure starting material.	materials. More tolerant to
		possible contaminants and
		less sophisticated extraction
		methods for preparation of
		starting material.
Sensitivity	Typically requires more than	Requires less than 10
	1000 polynucleotides/ml	polynucleotides/ml analyte.
	analyte.	
Specificity	Rate of false positives in	Rate of false positives in
	diagnostic kits can exceed	diagnostic kits can be much
	15%.	less than 12.5%. Improved
		specificity decreases need for
		costly post-processing and
		bioinformatics steps used in
		confirming the sequence of
		the target DNA or RNA.

The common denominator of PCR and PCR-based technologies practiced to date has been the use of thermal cycling to sequentially denature DNA, anneal primers, and then extend primers via a polymerase enzyme. The methods described herein, dubbed Nano-PCRTM, and apparatuses for performing those methods utilize the application of controlled amounts of force or stress to the nucleic acid molecules

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to provide new alternatives to thermal cycling for implementing DNA or RNA amplification. As used herein, applying stress to a nucleic acid includes direct and indirect application of force to a nucleic acid that tends to stretch or elongate the nucleic acid. As examples, stress can be applied to a nucleic acid by direct application of mechanical tension, by hydrodynamic stresses in a fluid flow, or electromagnetic fields, whether acting on the nucleic acid molecules themselves and/or on surfaces, substrates, or particles and the like that are bound to the nucleic acid. In many applications, Nano-PCRTM can break through one or more of the limitations that have traditionally restricted the performance and scope of conventional PCR.

Cycling of Mechanical Tension

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The application of controlled tension to nucleic acids provides not only an alternative to thermal denaturation of double-stranded DNA (dsDNA) but also a unique capability to precisely control each step of the PCR process. Nano-PCRTM introduces a new approach to amplification of DNA or RNA by exploiting the effects of precisely controlled forces, such as mechanical, hydrodynamic, or electromagnetic stresses on the DNA/RNA molecule and/or on the polymerizing enzyme.

Increasing temperature of solution comprising a DNA molecule and increasing stress on a DNA molecule produce analogous results. Thus, a polymerase reaction cycle can be initiated by increasing tension applied to a DNA template to above about 65 pN to denature the DNA. A step corresponding to the annealing of primers can be effected by slowly decreasing tension on the DNA template to below about 50 pN to allow primers to anneal to the template. Tension in the DNA template can then be adjusted within about 0 to about 30 pN during extension of the primer via enzymatic polymerization in order to control the progress, rate and/or accuracy of the replication. As with thermal cycling in conventional PCR, cycling of stress can be repeated in a pre-programmed cycle in Nano-PCRTM methods.

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In the examples below, various modes by which these methods can be put into practice are described. Hydrodynamic stresses and/or electric fields applied to the nucleic acid molecules, like mechanical tension can be cycled to perform Nano-PCRTM without reliance on thermal cycling. Of course, although Nano-PCRTM methods can be performed without any temperature cycling, this is not to say that control of temperature will not be advantageous in some embodiments as more fully discussed below. Figures 1A and 1B show exemplary flow diagrams for Nano-PCRTM methods. It will be appreciated that alterations and additions to the basic protocol will be made as appropriate for specific tasks such as sequencing, cloning, mutagenesis, mutation screening, and pathogen detection, etc.

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At room temperature and standard buffer conditions, application of tension above about 65 pN to double-stranded DNA can cause denaturation (i.e. melting) into single stranded DNA (ssDNA). As used herein, "room temperature" is understood to be a temperature within the normal range of comfortable laboratory temperatures, generally about 20-22° C. A theoretical model of the force- induced melting of DNA at room temperature has been described by Ioulia Rouzina and Victor A. Bloomfield ("Force-Induced Melting of the DNA Double Helix 1. Thermodynamic Analysis" *Biophys J.*, 80:882-93, 2001; and, "Force-Induced Melting of the DNA Double Helix. 2. Effect of Solution Conditions" *Biophys J.*, 80:894-900, 2001). Using the equations of Rouzina and Bloomfield as described herein, it is possible for one of ordinary skill to determine a precise level of tension that will melt a primer in a manner analogous to conventional melting point temperature calculations.

Slowly decreasing the applied tension below about 65 pN in the presence of primer oligonucleotides can permit the selective binding of the primers to template DNA in a manner analogous to slowly cooling denatured DNA below the melting point temperature of a primer in a thermal cycler. Accordingly, during a primer annealing step, tension applied to a nucleic acid template strand can be slowly reduced from an amount that causes dsDNA to melt to an amount that permits primer annealing. It may also be desirable to maintain tension on a DNA strand at a level that substantially inhibits polymerase action, for example at about 30 pN or

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greater but below about 50 pN, until unbound primer is flushed from the reaction chamber to minimize non-specific binding and non-specific primer extension.

The application of tension to a nucleic acid template in the range from about 0 to about 30 or 35 pN can be used to slow the rate of polymerase activity. The exact speed of the enzyme depends on various factors, including the ambient temperature or ambient concentrations of polymerase and/or nucleotide triphosphate substrates. Tension greater than about 35-45 pN at room temperature promotes the natural proofreading exonuclease activity of the polymerase enzyme.

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An exemplary embodiment of a Nano-PCRTM method can comprise: (a) providing a sample of double-stranded DNA (dsDNA) containing a target sequence, one or more oligonucleotide primers, for example a pair of primers complementary to the 3' ends of the target sequence and its complement; at least four different nucleoside triphosphates (i.e. ATP, CTP, GTP, TTP); and a DNA polymerase; (b) denaturing the dsDNA into single-stranded DNA (ssDNA) template strands using a non-thermally-driven process, for example by the application of tension sufficient to cause dsDNA to melt (*e.g.* tension greater than about 65 pN) to the dsDNA; (c) controlling the non-thermally-driven process to promote hybridization of primers to complementary template strands, for example, where tension was used to denature the dsDNA, by reducing the tension applied to the ssDNA; (d) permitting the DNAp to extend the primers to form dsDNA; and, (e) repeating steps (b-d) until a desired amount of DNA sequence amplification is obtained.

The use of a "non-thermally driven process" in the methods described herein means, for example, that dsDNA denaturation is not accomplished solely through an increase in temperature above the melting temperature of dsDNA, but rather that a physical or mechanical force is exerted on the nucleic acid that does not rely on temperature. The non-thermally driven process may comprise applying tension to the DNA strand, for example by direct application of mechanical force, by fluid flow, by application of an electric field, and/or by the action of one or more denaturing agents. As described herein, the effect of such a force may be affected by temperature so that it may be desirable in a given circumstance to control and optionally to modulate the temperature during one or more steps of the methods.

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A target sequence to be amplified can be contained in isolated DNA or in a mixture of nucleic acids and can be contained on complementary strands of equal or unequal lengths. A method may also include starting with a composition comprising RNA and producing a DNA template using reverse transcriptase or a similar method. A target sequence may alternatively be provided on a single stranded nucleic acid, rendering step (b) unnecessary in the first cycle. The reaction components of step (a) can be combined at the start of the procedure or may be introduced separately as needed. Optionally, reaction components can also be removed from the reaction chamber during certain steps. For example, nucleoside triphosphates (NTPs) may be introduced during or prior to step (d) and the primers may be introduced during or prior to step (c) and unbound primers may be flushed from the chamber before the primer extension (replication) step. Further, in various embodiments of the method, tension in the range of about 0-45 pN, about 0-35 pN, or about 0 to 20 pN can be applied to template DNA strands during step (d). In such embodiments, the amount of tension applied to the template strands during step (d) can optionally be varied over time. The amount of tension applied to the template strands in step (d) can varied according to the known or estimated progress of the polymerase in relation to positions of difficult or error prone subsequences, such as G-C rich segments (e.g. segments containing greater than about 50% G-C base pairs, or greater than about 70% G-C base pairs) or the positions of segments containing repeating sequences.

Accuracy of Nano-PCRTM over Normal and "Difficult" Sequences

When tension is applied to the template DNA during the primer extension step, a polymerase can be induced to "reverse direction" and the exonuclease activity of the polymerase can predominate. It will be appreciated that at the atomic scale and over times on the order of a single polymerase/exonuclease step, the process is stochastic. However, when considered from an average over the time scale of several steps the polymerase is seen to exhibit sustained exonuclease activity when the applied tension is greater than a threshold that can be theoretically predicted for a given temperature and solution conditions.

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By applying a modulated amount of tension to template DNA during the primer extension step in an amount below the threshold at which the exonuclease activity of a polymerase becomes predominant, e.g, below about 35 to 45 pN at room temperature and normal PCR solution conditions, more preferably in the range of about 10 to 30 pN, Nano-PCRTM can provide substantially increased accuracy of replicating DNA over conventional PCR. Furthermore, this effect can be achieved over those sub-sequences that are difficult using conventional PCR methods. The amount of tension applied to a template DNA strand can be adjusted in the range of about 0 to 45 pN, about 0 to 35 pN, or about 0 to 20 pN over time during the primer extension off a template that contains a mixture of more and less problematic segments. For example, according to a map of the sequence, tension may be increased as necessary to a level below about 35 pN to promote increased accuracy over difficult regions and then be carefully decreased to permit faster processing over less problematic segments. The length of the template strand is changed during the primer extension. In some variations of the methods, it is possible to adjust the tension on the template in direct response to the changes in length of the template strand so as to calibrate the applied tension precisely according to the progress of the polymerization reaction and the particular location of the "difficult" subsequences. In embodiments where it is not practical to directly monitor the progression of the polymerase, the position of the polymerase can be estimated by multiplying the elapsed time by the known rate of replication for the polymerase at the applied amount of tension.

Thus Nano-PCRTM permits accurate replication of not only normal target templates but also difficult sequences (e.g. GC rich DNA, tandem repeat, microsatellite or trinucleotide repeat DNA) to be replicated and amplified with substantially increased accuracy relative to conventional PCR. As an example, where one of the highest fidelity polymerases currently available (e.g. Phusion Enzyme) is used in conventional thermally-driven PCR, error rates of approximately 4.0×10^{-7} errors/base pair are observed in favorable cases, that is, only on well-behaved sequences. By contrast Nano-PCRTM methods as described herein can produce an error rate less than about 1.0×10^{-7} errors/base pair, less than 5.0×10^{-8}

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errors/base pair, 1.0×10^{-8} errors/base pair, 5.0×10^{-9} errors/base pair, 1.0×10^{-9} errors/base pair, 5.0×10^{-10} errors/base pair, or even 1.0×10^{-10} errors/base pair. Furthermore the methods described herein can permit the efficient amplification of oligonucleotide fragments having a GC content higher than 50 percent, 60 percent, 70 percent, 75 percent., or even 80 percent or 85 percent.

In certain cases, oligonucleotide fragments contain a section of repeating base pair units at least eight base pairs in length (*e.g.*, AAAAAAA, GCGCGCGC). The error rate for conventional PCR is increased in such cases, usually approaching 1.0×10^{-2} errors/base pair. The present methods of Nano-PCRTM provide for the amplification of repeating base pair units with an error rate less than about 1.0×10^{-3} errors/base pair. Under certain conditions, error rates less than 1.0×10^{-4} errors/base pair, 1.0×10^{-5} errors/base pair, 1.0×10^{-6} errors/base pair, or 1.0×10^{-7} errors/base pair can be achieved. These low error rates may also be obtained where the repeating base pair unit is at least 10 base pairs in length, at least 15 base pairs in length, or at least 20 base pairs in length. In the methods described herein, such results can also be obtained over microsatellite regions, polymerase slippage regions, and other tandem repeat regions, which are difficult sequences when using conventional PCR methods.

20 Amplification Efficiency

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Where amplification efficiency is defined by the equation $N_1 = N_2(1+Y)^n$, and N_1 is the number of product copies, N_2 is the number of template oligonucleotide copies, n is the number of cycles and Y is the efficiency, efficiencies greater than 80 percent are achieved. Under certain conditions, amplification efficiencies greater than 85 percent, 90 percent, 95 percent, 96 percent, 97 percent, 98 percent, or even 99 percent can be achieved. Such amplification efficiencies can also be obtained where the GC content of the oligonucleotide fragment is greater than 55 percent, 60 percent, 65 percent, 70 percent or even 75 percent.

Amplification efficiencies greater than 50 percent can be achieved where the GC content of an oligonucleotide fragment is greater than 50 percent, 55 percent, 60 percent, 65 percent or 70 percent. Efficiencies greater than 60 percent, 70 percent,

80 percent, 90 percent, 95 percent, 97 percent, or even 99 percent can be observed. In conventional PCR, these high GC-rich regions are amplified and sequenced with the addition of various denaturing agents, including but not limited to sodium hydroxide, TMA chloride, TMA oxalate, TMA acetate, TMA hydrogen sulfate, ammonium chloride, benzyldimethylhexadecylammonium chloride, HTA bromide, HTA oxalate, betaine monohydrate, DMSO, and formamide, and the like. In certain embodiments of Nano-PCRTM, efficient amplification is also seen using methods as described herein in the absence of such polymerase chain reaction additives.

10 Robustness and Adaptability

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Conventional PCR methods generally rely on precise control and cycling of temperature. Further, conventional PCR methods can require additional factors such as various denaturing agents and tedious optimizations. However, the use of tension cycling to drive amplification as described herein enables a much higher degree of precision and control over the PCR process than allowed by thermal cycling alone. Moreover, the methods described herein can function under a wide range of temperature conditions, limited only by factors such as the range of temperatures under which a chosen polymerase can function and the melting point temperature of the primer/template bond at a given tension.

Of course, it will be appreciated that temperature can affect the rate and accuracy of polymerase enzymes and the melting of DNA under tension. Generally, the amount of tension required to melt dsDNA is decreased with increasing temperature. As a rough guide, from about to 0 to 20° C, up to about 75 pN can be required to melt dsDNA. At about 60° C, the amount of tension required to denature dsDNA can be about 45 pN. The melting tension decreases to about 7 pN at just below the free DNA melting point.

Although it is generally not necessary, it may be advantageous to control the temperature during one or more steps of the methods depending on requirements of an individual application. For example, the temperature of the reaction mixture can be maintained at a temperature that optimizes the accuracy, polymerization rate, and/or tension response of a chosen DNA polymerase, that increases or decreases

the amount of tension required to achieve DNA melting, or that is otherwise advantageous because of the individual device or working environment. Unless otherwise desired, the temperature can be generally constant, and the entire process can be performed at or near normal room temperature. Unless otherwise indicated, the examples herein are described using amounts of tension that will be appropriate at room temperature. One of ordinary skill will readily be able to adjust the tension applied to the DNA template for higher or lower temperatures.

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Through the application of even small amounts of tension, thermal cycling temperatures no longer impose a limitation on the temperature at which the PCR reaction must be carried out. By applying a tension of about 7 to 45 pN, for example, one can decrease the temperature at which double-stranded DNA denatures by up to about 30 degrees C. Adjusting the amount of tension applied to DNA enables performance of PCR at temperatures well below the amount required for denaturation in conventional PCR. This effect permits PCR using low amplitude thermal cycling. For example, a method can comprise a denaturation step in which an amount of force less than about 65 pN is applied in concert with an increase in the temperature of the solution to less than about 90° C, alternatively less than about 80° C.

The methods and apparatuses described herein may be carried out or operated at temperatures below 90° C. Oligonucleotide denaturation steps, for instance, can be conducted at below 80° C, 70° C, 60° C, 50° C, 40° C, 30° C, or even 25° C. Annealing steps can be conducted below 50° C, 45° C, 40° C, 35° C, 30° C, or even 25° C. Furthermore, polymerization steps can be conducted below 70° C, 60° C, 50° C, 40° C, 30° C, or even 25° C.

Similarly, pH and ionic strength of the solution in which the DNA is immersed can affect the tension-induced melting curves of DNA. Accordingly, by adjusting the levels of force applied to DNA in the methods, Nano-PCRTM permits using a wider range of pH and ionic strength solution conditions to carry out the PCR process than in conventional PCR. Similarly, all these and additional parameters can affect tension control of primer extension. The methods described herein can be adjusted for and even take advantage of these effects. Thus Nano-

PCRTM methods can be more robust to a wider range of temperature, ionic strengths, pH and buffer conditions in general, which means that it can be performed in a wider range of situations, demanding less stringent extraction and purification of the starting DNA or RNA material, and can be more resistant to various contaminants and enzyme inhibitors that typically restrict the scope of conventional PCR. In preferred implementations, the presence of contaminating substances can be removed by flushing the sample as part of the Nano-PCRTM process. For example, an unpurified DNA sample containing contaminants can be introduced into a Nano-PCRTM device, the DNA is retained in the reaction chamber, *e.g.* by any of the means described herein for retaining DNA for the controlled application of stress. The contaminants are flushed out of the reaction chamber and reagents are flushed in. This can provide substantial robustness to the Nano-PCRTM process, permitting rapid accuracte amplification in environments that are unfavorable to conventional PCR.

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$Nano-PCR^{ ext{ iny }}M$ Using Direct Application of Mechanical Force

Nano-PCRTM methods can be performed utilizing various methods to directly apply mechanical tension to DNA strands as a non-thermally-driven process that can provide for DNA denaturation and/or precise control of the activity of DNA polymerase. There are several different ways to apply tension to a double-stranded oligonucleotide. For example, DNA strands may be anchored in an array to a movable element, to individually controllable elements, or to particles that can be manipulated.

25 Using opposed coated surfaces:

As an example, the process can be performed using nucleic acids anchored to opposed coated surfaces, generally as illustrated in Figures 2A-2C: Coated surfaces are prepared by attaching a first complexing molecule (e.g., streptavidin) 201, 207, which can be the same or different for each surface, to two substrate surfaces 203, 209. The coated substrate surfaces are arranged in opposition to one another at a suitable distance apart. Double-stranded nucleic acids 205, where both ends of one

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strand comprise a complexing molecule that is complementary to the first complexing molecule (*e.g.*, biotin) which can recognize and bind to the first complexing molecule, are immobilized onto the coated surfaces. Force can be applied to the ends of the immobilized nucleic acids by increasing the distance between the coated surfaces or by lateral translation of one or both surfaces. For example, one or both substrates may be a movable element or comprise a movable element, such as a piezoelectic element. Tension sufficient to cause dsDNA to melt (*e.g.* greater than about 65 pN at room temperature) can be applied to the nucleic acids, producing anchored strands and freed strands. Both the anchored and the freed strands can be replicated using appropriate primers and polymerase. Preferably, the freed strands can be flushed away and optionally collected so that only the anchored strands will be replicated. The position of the opposed surfaces can be controlled during replication to modulate the amount of tension applied to the anchored template strands. The cycle can be repeated as desired.

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In variations of a device for performing the method using opposed coated surfaces 215, 217, one or both surfaces can also be arranged to form an array of individually movable elements 219, each of which may be individually addressed by a control circuit driven by a programmable processor as illustrated in Figure 2C. Such a control circuit can include a feedback channel that reports force and/or displacement parameters to the processor. Printing or lithography techniques can be used to pattern sites for anchoring molecules on a surface. A device for performing the method can also comprise a channel for introducing reagents to the chamber or channel comprising the coated surfaces and apparatus for delivering (and optionally storing) reagents separately or in combination and for collecting reaction products. A wide variety of suitable methods of anchoring a nucleic acid to a surface are known, including but not limited to covalent bonding, antigen-antibody, and streptavidin-biotin.

Arrangements of fluid flow can be utilized to orient and extend DNA strands between opposed surfaces. For example, as illustrated in Figure 2B, DNA may be anchored at one end to a surface 213 having passages for fluid flow distributed between the anchoring locations. Flowing fluid though these passages can be used

to orient and extend DNA strands more or less uniformly in a desired direction, for example towards an opposed surface 211 or array of movable elements, which may have passages distributed between anchoring surfaces to receive the fluid flow. Thus, a method may comprise anchoring DNA strands to a first surface, flowing fluid through openings in the first surface towards and through openings in a second surface opposed to the first surface, and anchoring DNA strands oriented in the fluid flow to the second surface.

To increase the number of anchored strands with each cycle, activatable primers can be used in replicating the attached strands. "Activatable primers" comprise chemical moieties that can be activated by chemical or physical methods. These "activatable groups" are inert until activated, for example by photoactivation using a laser at an appropriate wavelength. Many different activatable chemical groups are known in the art, which can be converted into or unblock functional complexing groups. In a variation of the method, activatable primers are allowed to anneal to the anchored single-stranded nucleic acids. Primer extension and fragment replication is performed. The resulting double-stranded nucleic acids are denatured through the application of tension to the template strand. The free copy strands will then comprise the activatable groups of the primers. Activation allows the copy strands to become immobilized on the opposing coating surfaces. This cycle can be repeated until a desired degree of amplification is obtained. When desired, anchored nucleic acids can be released, for example by the use of a restriction enzyme that recognizes a sequence near an anchored end of the nucleic acid or that has been introduced into the end of the copied nucleic acid by the primers.

25 Using optical or magnetic traps:

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Another way to directly apply tension to DNA can utilize optical or magnetic tweezers or other traps to manipulate particles to which the DNA is anchored. An optical tweezers traps particles with forces generated by optical intensity gradients. Dielectric particles polarized by the light's electric field are drawn up the gradients to the brightest point. Reflecting, absorbing and low-dielectric particles, by contrast, are driven by radiation pressure to the darkest point. Optically generated forces

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strong enough to form a three-dimensional trap can be obtained by bringing a laser beam with an appropriately shaped wavefront to a tight focus with a high numerical aperture lens. Figure 3A illustrates a DNA strand extended between beads 301 tapped at the focus of a laser beam 303. It is possible to manipulate large numbers of particles using an array 307 of optical tweezers as illustrated in Figure 3B. Commercially available optical tweezers arrays include those produced by Arryx, Inc. Another implementation of an array of optical tweezers, see E. R. Dufresne and D. G. Grier, Rev. Sci. Instr. 69:1974 (1998); and, U. S. Patent No. 6,055,106 (2000). An optical tweezers array can comprise about 10^3 , 10^4 , 10^5 , 10^6 , or more pairs of optical or magnetic tweezers.

Amplification using optical or magnetic tweezers can generally be performed as follows: A nucleic acid is anchored to appropriate particles at each end in a fluid medium. The particles may be adapted to be manipulated using optical or magnetic traps. Tension sufficient to denature a dsDNA, for example greater than about 65 pN, is applied to the oligonucleotide through the application of force (e.g., optical or magnetic) to the particles, resulting in the denaturation of the nucleic acid. The tension can be reduced in the presence of primers to allow the primers and nucleic acid to anneal. Polymerization by DNA polymerase can be initiated by further relaxing the tension. To repeat the cycle, tension can be increased such that the resulting double-stranded nucleic acids are denatured. In a variation, a nucleic acid can be anchored at one end to a bead that is trapped in a fluid flow, for example by a magnetic field. Fluid flow rate can be used to control tension on the nucleic acid.

It is possible to begin from a single target molecule and sequentially populate an array with copy strands. Copy strands can be anchored to new beads using activatable primers. New beads can be brought into proximity with the copied strands. Alternatively, beads having pre-immobilized primers can be brought into proximity with the copied strands in conjunction with the denaturation step. Manipulation of the beads in this fashion optionally may be automatically controlled by a programmable processor.

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Nano-PCRTM Using Hydrodynamic Stress

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Nano-PCRTM methods can be performed utilizing the application of tension to DNA by hydrodynamic stress in controlled fluid flow. Methods using this approach can be performed in a microfluidic device, which can be a benchtop device or alternatively can be reduced to a portable size such as may be incorporated in a handheld device. Nano-PCRTM methods utilizing the application of tension to DNA by hydrodynamic stress in controlled fluid flow can be performed using any arrangement that provides for a controlled rate of fluid flow.

Using anchored DNA polymerase: A method of performing a Nano-PCRTM method using polymerase anchored to a surface in a device can comprise the following steps. Polymerase is immobilized on a surface that is arranged such that fluid can be flowed over the surface at a controlled rate. For example, a surface in a channel or chamber that has been coated with a first complexing moiety can be used to immobilize a DNA polymerase that has been modified to comprise a second complexing moiety. Exemplary complexing moieties include antigen-antibody, histidine to Ni-NTA, or biotin- streptavidin pairs. Target dsDNA is denatured in the presence of primers, for example dsDNA and primers can be subjected to a flow rate such that a force sufficient to cause dsDNA to melt (e.g. greater than about 65 pN) is applied to the dsDNA. Polymerase/nucleotide/primer complexes can be allowed to form by reducing the flow rate. Primer extension and fragment replication can be promoted through a further reduction of flow rate. Double-stranded nucleic acid products comprising template and copy strands can be denatured through the application of an increased flow rate, and the cycle can be repeated until a desired degree of amplification is obtained. In such a method, polymerase may be immobilized in a microchannel, for example a microchannel in a microfluidic "lab on a chip" type device.

In variations of a device for performing the method using polymerase anchored on one or more coated surfaces, such a device can also comprise a channel for introducing reagents to a reaction chamber or channel comprising the coated surfaces and apparatus for delivering and optionally storing reagents separately or in combination and for collecting reaction products.

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Printing or lithography techniques known in the art can be used to pattern sites for anchoring molecules. Such a device will comprise an apparatus for creating and controlling fluid flow in the reaction chamber or channel. Any suitable method for creating and controlling fluid flow can be used including electrodynamic methods, pumps and syringe apparatuses. Reagent solutions optionally can be recycled through the reaction chamber.

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Figure 4 illustrates an approach in which polymerase 401 is anchored to a substrate 407, for example by streptavidin binding and the like. DNA strands 403 are permitted to bind to the anchored polymerase. Controlled fluid flow 409 passed over the substrate 407 causes application of stretching force on the DNA strands in the form of hydrodynamic stress. Nano-PCR can be carried out using such an application of force according to the general scheme illustrated in Figure 1 as described above.

Using anchored DNA strands in a controlled fluid flow: Another way to apply tension to nucleic acids involves immobilized nucleic acids in a controlled fluid flow. The process can generally be performed as follows: Nucleic acids comprising, at one end, a first complexing moiety that recognizes and can bind to a second complexing moiety coated on a surface, are allowed to become immobilized on the coated surface. Fluid is flowed over the surface such that a force sufficient to cause dsDNA to melt (e.g. greater than about 65 pN) is applied to anchored doublestranded nucleic acids, which results in strand separation. DNA polymerase and primers, optionally primers comprising activatable groups, are flushed over the surface at a reduced flow rate. A reduced or stopped flow rate allows formation of polymerase/oligonucleotide/primer complexes. Primer extension and fragment replication can be promoted in the presence of NTPs through a further reduction or stoppage of flow rate. After replication, the flow rate can be increased, subjecting the resulting dsDNA to tension such that the dsDNA is denatured. If activatable primers are used, the extended primers can be activated. These activated, extended primers can be allowed to bind to the coated surface, and the cycle can repeated until a desired degree of amplification is obtained. In such a method, polymerase

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may be immobilized on a surface in a microchannel, for example a microchannel in a microfluidic "lab on a chip" type device.

Figure 5 illustrates the stretching of DNA in a fluid flow by hydrodynamic stress where DNA strands 503 are anchored to a substrate 507 through anchoring molecules 501. Fluid flowing in direction 509 extends and stretches the DNA in a controlled manner as a function of the fluid flow velocity. Figure 5B illustrates a variation in which DNA strands 503 are anchored by binding molecules 501 to a plurality of substrate structures 505 such that fluid can flow between the structures at a controlled rate. It will be appreciated that there are a large number of other variations that can be used to achieve a similar result.

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In variations of a device for performing the method using nucleic acid anchored on one or more coated surfaces, such a device can also comprise a channel for introducing reagents to a reaction chamber or channel comprising the coated surfaces and apparatus for delivering and optionally storing reagents separately or in combination and for collecting reaction products. Printing or lithography techniques known in the art can be used to pattern sites for anchoring molecules. Such a device can comprise an apparatus for creating and controlling fluid flow in the reaction chamber or channel. Any suitable method for creating and controlling fluid flow can be used. For example flow can be provided by means of a pump or can be electrostatically driven. Reagent solutions can optionally be recycled through the reaction chamber.

Stretching DNA in a velocity gradient and using hydrodynamic focusing: An alternative approach to applying tension using fluid flow can be used in combination with the above methods, or may form the basis of a distinct method. DNA can be stretched in a fluid that has a velocity gradient. Various exemplary arrangements for hydrodynamic focusing and counter propagating elongational flows are illustrated in Figures 6A-6C.

For example, Wong et al. reviewed the basis of several such techniques and described a method of hydrodynamic focusing (Wong et al., "Deformation of DNA molecules by hydrodynamic focusing." J. Fluid Mech. 497:55-65, 2003). In hydrodynamic focusing, illustrated by Figure 6A, two streams of buffer 607 flowing

at a relatively high rate converge in a microchannel 605 with a center stream that is introduced at a low flow rate. The converging streams accelerate the center stream without substantially mixing. The result is a region of flow having a strong velocity gradient in the flow direction. DNA 601 in this gradient is stretched to an extended state. By increasing the flow rates of the converging streams even more, it will be possible to denature dsDNA such that ssDNA emerges from the microchannel. This permits delivering ssDNA to a reaction chamber, for example where polymerase has been anchored.

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Stagnation flow can be used to trap and apply tension to nucleic acids without the need for any anchoring. Perkins et al. described elongation of DNA in a planar elongation flow apparatus ("Single Polymer Dynamics in an Elongation Flow" Science, 276:2016-21, 1997). In Perkins' apparatus, fluid is flowed 623 from opposing directions into a T-shaped junction 625 such as illustrated in Figure 6C. At the center of the junction, a stagnation point 629 is established. Outside of this point, a fluid velocity gradient is established. DNA 601 can become trapped at the stagnation point, being pulled equally in opposite directions by the velocity gradient. Alternative arrangements such as channel 615 illustrated in Figure 6B can include offset jets 617 of fluid entering a channel 615, or flowing buffer in opposing directions across a slot in which nucleic acid resides.

20 Nano-PCRTM Methods Using Cycling of Applied Electric Fields

It is possible to apply force to DNA strands in Nano-PCRTM methods through the use of electric and magnetic fields. There are a variety of ways that this can be accomplished. For example, electric fields can be used to indirectly apply force to DNA by driving fluid flow in a microfluidic device. As described above, fluid flow can be used to apply hydrodynamic stress to DNA, for example, DNA anchored to a surface and/or to a particle, or bound to a DNA polymerase that is anchored to a surface. Electrophoretic forces can also apply force directly to DNA strands.

Electric fields may also be used to manipulate DNA strands bound to conductive particles. Accordingly, Nano-PCRTM methods can be performed where denaturation, annealing, and/or primer extension steps are controlled by a non-

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thermally-driven process wherein one or both ends of a DNA strand is bound to a conductive particle, *e.g.* gold nanoparticles or the like, which can be manipulated by electric fields to apply tension to the DNA strand. Where one end of a DNA molecule is attached to a conductive particle, the other end can be anchored to a surface in a reaction chamber in a device. Such methods may utilize activatable primers as described herein to anchor DNA strands produced in each cycle.

Exemplary Applications of Nano-PCRTM Methods

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Nano-PCRTM methods can be employed in kits and systems for pathogen and bioweapon detection. Examples of such pathogens include, without limitation:

Adeno-associated Virus (AAV), Adenovirus, Cytomegalovirus (CMV), Epstein-Barr Virus, Enterovirus, Hepatitis A Virus (HAV), Hepatitis B Virus (HBV),

Hepatitis C Virus (HCV), Human Herpes Virus Type 6 (HHV-6), Human Immunodeficiency Virus Type 1 (HIV-1), Human Immunodeficiency Virus Type 2

(HIV-2), Herpes Simplex Virus Type 1 and Type 2 (HSV-1 and HSV-2), Human T-Cell Lymphotropic Virus Type I and Type II (HTLV-I and HTLV-II),

Mycobacterium tuberculosis, Mycoplasma, Parvovirus B-19, Respiratory Synctitial Virus (RSV) and Porcine Endogenous Retrovirus (PERV). Nano-PCRTM methods can be used for detection of any pathogen in any environment because of the enhancements in sensitivity, accuracy and robustness these methods can provide.

The detection and identification of a particular pathogen using conventional PCR-based diagnostics generally requires that the pathogenic organism or its polynucleotide be present in a biological fluid (e.g. blood, saliva, etc.) at a certain threshold concentration. The lower detection limit of *Mycobacterium tuberculosis*, for example, has been reported as 7.5 x 10³ organisms/ml. HCV RNA is detectable in a range from 100 to 1000 RNA molecules/ml. Shim has reported that the polymerase chain reaction detects around 87.5 percent of proven *Mycobacterium tuberculosis*-containing nodules. That corresponds to a false-negative rate for detection of 12.5 percent. In preferred embodiments Nano-PCR™ methods can be used to detect pathogens such as the above with false-negative rates typically less

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than 12.5 percent, 10 percent, 5 percent, 2.5 percent, 1 percent, 0.5 percent, 0.25 percent, or 0.1 percent.

Furthermore, Nano-PCRTM, can be performed such that it is not limited to about 30-35 cycles of amplification as conventional PCR generally is. This is because of degradation of polymerase after repeated cycles of heating above the DNA melting temperature. In contrast, Nano-PCRTM methods can optionally comprise 40, 50, 60, 70, 100, or more cycles. As Nano-PCRTM can be performed in an isothermal manner, or using low amplitude temperature modulation, Nano-PCRTM can be repeated for many cycles, limited only by the lifetime of the enzyme (e.g. at room temperature).

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Thus Nano-PCRTM methods can be used to amplify amounts of starting material (either organisms or their DNA or RNA) that are substantially less than amounts required by conventional PCR. Nano-PCRTM methods can be used to detect and reliably amplify as little as a single molecule of DNA or RNA, dramatically decreasing the false-negative rate and providing increased sensitivity of as much as 100%. For pathogens such as those exemplified above, organisms or polynucleotides can be detected at concentrations lower than 1000 organisms or polynucleotides/ml, 100 organisms or polynucleotides/ml, 50 organisms or polynucleotides/ml, 25 organisms or polynucleotides/ml, 10 organisms or polynucleotides/ml, 5 organisms or polynucleotides/ml, or even as little as 1 organism or polynucleotide/ ml.

An exemplary variation of the method can be used for detecting the presence or absence of at least one specific DNA sequence or distinguishing between two different DNA sequences in a sample. In such a variation, target DNA can be amplified as described above. The method can further comprise: contacting the amplified DNA with a probe or probes (*e.g.*, an oligonucleotide complementary to the sequence to be detected that also comprises a detectable moiety, such as a fluorescent label); and, detecting whether the specific DNA sequence is in the sample by observing the presence or absence of the probe bound to the amplified DNA, or distinguishing between two different sequences by detecting which of a plurality of probes is bound to the amplified DNA.

Another variation of the method can be used for amplification and/or detection of a sequence encoded on RNA. The target sequence can be encoded on an isolated RNA or on RNA in a mixture of nucleic acids. The method can comprise: isolating RNA from a sample (e.g., tissue or fluid); performing reverse transcription thereby obtaining a corresponding cDNA; and, amplifying the target sequence as described above. Such methods can further comprise detecting the presence of a specific sequence in the sample as described above.

Another variation of the method can be used for sequencing a DNA. Such a method can comprise optionally amplifying the DNA as described above and sequencing the DNA. Sequencing the DNA can comprise (a) providing a sample of dsDNA containing a target sequence, the sample being divided into four parallel reactions, a primer complementary to the 3' end of the target sequence; at least four different nucleoside triphosphates (i.e. ATP, CTP, GTP, TTP); providing a different dideoxy nucleoside triphosphate (ddNTP) selected from among ddATP, ddCTP, ddGTP, and ddTTP optionally labeled with a detectable chemical moiety such as a fluorescent moiety, and a DNA polymerase in each parallel reaction; (b) denaturing the dsDNA into ssDNA template strands using a non-thermally-driven process, for example by the application of tension sufficient to cause dsDNA to melt (e.g. greater than about 65 pN) to the dsDNA; (c) controlling the non-thermally-driven process to promote hybridization of primers to complementary template strands, for example, where tension was used to denature the dsDNA, by reducing the tension applied to the ssDNA; (d) permitting the DNAp to extend the primers to form dsDNA; (e) optionally repeating steps (b-d) until a desired amount of DNA sequence amplification is obtained, and determining the sequence by detecting the length of each nucleotide produced in the reaction or by detecting the base specific fluorescent moiety or some other base-specific signal as in various single molecule sequencing schemes.

Nano-PCRTM Devices

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There are many different device types and configurations one can use to perform non-thermally-driven polymerase chain reactions as described herein. One

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such device is a microfluidic device, where the flow rate within microfluidic channels on the device is controllable and variable. In preferred embodiments, a device will have a reaction chamber, which can be a channel, an arrangement of channels, or an enclosed space. The reaction chamber will generally comprise a means of retaining nucleic acids and a means of applying stress or tension to the nucleic acids retained therein. Thus, arrangements designed to carry out any of the methods described herein can be envisioned comprising a combination of channels and enclosed spaces having disposed therein particles capable of binding nucleic acids or complexing molecules capable of securing their complementary complexing molecules, surfaces having complexing molecules, movable elements, channels for directing fluid flow and generating a fluid velocity gradient, pumps, valves, membranes, and the like. The chamber can comprise an optically transparent window, for example, if optical micromanipulators are to be used. The devices can be manufactured as microfluidic devices which may be incorporated into handheld units. If desired, Nano-PCRTM can be performed in solution volumes of less than about a microliter, for example about 50-1000 nL, preferably about 100-500 nL.

As an example, Figure 7A illustrates a possible configuration of a device in which reagents can be introduced through inlets 701, 707 and 715. One or more storage chambers 705, 711 can be provided to contain prepared buffers, dideoxy nucleotide triphosphates, polymerase, and the like. Valves 703, 709, 717 and 718 may comprise one or more fluid gates arranged to control fluid flow at junctions between channels. Reaction chamber 715 may be arranged to permit controlled application of tension to nucleic acid molecules therein, for example as illustrated in Figures 2-6. A channel 721 and pump 723 are optionally provided to permit recycling and controlled flow of regents through chamber 715. Pump 723 may operate by any appropriate mechanism recognized in the art, for example peristaltic pumping, pumping by use of one or more bellows or pistons, by electromotive force, and variations or combinations of such devices and the like. Where recycling is not desired, flow may be controlled within chamber 715 or externally, for example by syringes attached at inlet and/or outlets 701 and 719. An example of a microfluidic device utilizing a circular, or roughly circular, channel configuration is illustrated by

Figure 7B. Inlets 751 permit introduction of reagents either directly to a channel feeding reaction channel 763 or into one or more storage chambers 753, 754 for later use. Valves 755, 757, and 759 control flow into and out of the reaction channel. Pumps 761 may operate to control fluid velocity in channel 763 by peristaltic action, for example by deflection of one or more valve gates into channel 763 in a sequentially controlled manner, electromotive force, or any other means recognized in the microfluidics art. For example, a device may be constructed using valves and a peristaltic pumping arrangement that comprise structures constructed of elastomeric material that can be deflected into the channels of the device in a controlled sequence to control flow, such as described in published PCT application WO/02081729.

The operation of the device illustrated in Figure 7B can be further understood through a description of its operation during a non-thermally-driven polymerase chain reaction. In the specific instance of nucleic acid amplification reactions, a sample containing or potentially containing a target nucleic acid is introduced into loop 763 through an inlet 751. In some examples, one or more walls of the loop 763 have been prepared for anchoring polymerase or nucleotides as illustrated in Figures 4-5. Alternatively, the loop may be arranged to create fluid velocity gradients, counter propagating fluid flow, and the like by utilizing additional inlets or rotating surfaces such as illustrated in Figure 6. Other reagents necessary to conduct the amplification reaction are similarly introduced through the inlets. Typical reagents include a primer or primers (e.g., forward and reverse primers) that specifically hybridize to the target nucleic acid, the four deoxynucleoside triphosphates (i.e., dATP, dTTP, dGTP and dCTP), a polymerase, a buffer and various cofactors required by the polymerase (e.g., metal ion).

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Following introduction of the sample and necessary amplification reagents into loop 763, the resulting solution is circulated under the action of pumps 761. By varying the rate of pump action, one can control the solution circulation/flow rate. A flow rate resulting in application of about 65 pN of force to the target nucleic acid is established, which denatures it. The flow rate is decreased such that the force applied to the target nucleic acid is in a range from about 30 pN to 60 pN. This

allows formation of polymerase/nucleic acid/primer complexes. Primer extension is initiated by further reducing the flow rate to a value corresponding to less than 30 pN of applied force. Upon completion of primer extension, the flow rate is again increased to denature the resulting double-stranded nucleic acid. The recited steps are repeated until a desired quantity of target nucleic acid is obtained. One can access the amplified target nucleic acid by flushing solution through outlet 765 by opening valve 759.

An apparatus for conducting Nano-PCRTM methods can comprise a programmable control device that can individually address and control elements of the reaction device and may also include sensors and feedback circuits so that the control device can monitor, analyze, and if desired can adjust reaction parameters, such as applied stress and template extension.

Examples

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Example 1: Method and Device Using Opposing Coated Surfaces

A pair of streptavidin-coated surfaces are prepared according to standard methods. (Sabanayagam, Smith, and Cantor. "Oligonucleotide immobilization on micropatterned streptavidin surfaces." Nucleic Acids Res. 2000, Vol. 28, No. 8 pp. i-iv) Biotinylated dsDNA (biotinylation at both ends of one strand) is added to the surfaces, which immobilizes the dsDNA between the surfaces. Jeffrey M.Rothenberg and Meir Wilchek. p-Diazobenzoyl-biocytin: a new biotinylating reagent for DNA Nucleic Acids Research Volume 16 Number 14 1988) By adjusting the concentration of the template that is applied to the surface, the surface density of the DNA molecules can be controlled. At room temperature, greater than about 65 pN of tension is applied to the dsDNA by increasing the distance between the coated surfaces. This denatures the dsDNA, leaving only target ssDNA for amplification. The bound DNA is contacted with primers comprising caged biotin groups, while the distance between the surfaces is reduced so that between 30 pN and 60 pN is applied to the immobilized ssDNA. The primers are allowed to anneal to the target DNA, and DNA polymerase and nucleotides are added to the resulting complex. Primer extension is initiated by further reducing the applied tension to

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<30 pN. Once primer extension is complete, a force >65 pN is applied to the resulting duplex by increasing the distance between the surfaces. This application of force denatures replica nucleotide strand from its template. The replica strands containing caged biotin moieties are photoactivated and allowed to bind to the streptavidin-coated, opposing surfaces. The above-recited steps are repeated until a desired degree of amplification is obtained for the target nucleotide.

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Caged biotin reagents can be purchased from commercial vendors such as Molecular Probes or Pierce. For example, a derivative of biotin with a photoactivatable nitrobenzyl group (MeNPOC-biotin) exists in a form well-suited for easy linkage to biomolecules and surfaces. (Pirrung MC, Huang CY. A general method for the spatially defined immobilization of biomolecules on glass surfaces using "caged" biotin. Bioconjug Chem. 1996 May-Jun;7(3):317-21).

Example 2: Method and Device Using Immobilized Polymerase

A streptavidin-coated microchannel surface is prepared according to standard methods. Sabanayagam, Smith, and Cantor. "Oligonucleotide immobilization on micropatterned streptavidin surfaces." Nucleic Acids Res. 2000, Vol. 28, No. 8 pp. i-iv) Biotinylated DNA polymerase is flushed into the microchannel and incubated to allow surface saturation. Commercial kits for the biotinylation of enzymes are available, for example, from Pierce Labs. Unbound enzyme is flushed out of the microchannel and target nucleotide (e.g., ssDNA, RNA) and primers are flushed in at a chamber flow rate that applies >60 pN of force on the nucleotide. The polymerase/nucleotide/primer complex is allowed to form by reducing the flow rate such that a force between 30 pN and 60 pN is applied to the nucleotide. Primer extension is allowed to occur by further reducing the chamber flow rate to <30 pN. Once primer extension is complete, a force >65 pN is applied to the resulting duplex by increasing flow rate. This application of force denatures replica nucleotide strand from its template. The denatured strands are allowed to cycle through the microfluidic chamber until polymerase binding occurs, and the above-recited steps are repeated until a desired degree of amplification is obtained for the target nucleotide.

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Example 3: Method and Device Using DNA Immobilization

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A streptavidin-coated microchannel surface is prepared according to standard methods. (Sabanayagam, Smith, and Cantor. "Oligonucleotide immobilization on micropatterned streptavidin surfaces." Nucleic Acids Res. 2000, Vol. 28, No. 8 pp. i-iv. Biotinylated dsDNA (biotinylation at one end of one strand) is flushed into the microchannel and incubated to allow surface binding. (Jeffrey M.Rothenberg and Meir Wilchek. p-Diazobenzoyl-biocytin: a new biotinylating reagent for DNA Nucleic Acids Research Volume 16 Number 14 1988) A chamber flow rate that applies a force >65 pN to the bound dsDNA is established. This denatures the dsDNA, leaving only target ssDNA for amplification. DNA polymerase, nucleotides, and caged biotinylated primers are flushed into the microchannel. Caged biotin reagents can be purchased from commercial vendors such as Molecular Probes or Pierce. For example, a derivative of biotin with a photoactivatable nitrobenzyl group (MeNPOC-biotin) exists in a form well-suited for easy linkage to biomolecules and surfaces. (Pirrung MC, Huang CY. A general method for the spatially defined immobilization of biomolecules on glass surfaces using "caged" biotin. Bioconjug Chem. 1996 May-Jun;7(3):317-21). The chamber flow rate is decreased such that between 30 pN and 60 pN is applied to the bound ss DNA. The primers are allowed to anneal to the target DNA, and the resulting compounds are allowed to complex to DNA polymerase. Primer extension is allowed to occur by further reducing the chamber flow rate to <30 pN. Once primer extension is complete, a force >65 pN is applied to the resulting duplex by increasing flow rate. This application of force denatures replica nucleotide strand from its template. The replica strands containing caged biotin moieties are photoactivated and allowed to bind to the streptavidin-coated microchannel surface. The above-recited steps are repeated until a desired degree of amplification is obtained for the target nucleotide.

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Example 4: Method and Device Using Optical Tweezers

A double-stranded DNA complex is immobilized between polystyrene beads in an appropriate medium at ambient temperature. ("Overstretching B-DNA: the Elastic Response of Individual Double Stranded and Single Stranded DNA Molecules" by Steven B. Smith, Yujia Cui, and Carlos Bustamante Science (1996) 5 vol. 271, pp. 795-799) A stretching force of approximately 65 pN is applied to the DNA through the use of optical tweezers. (Rouzina, I., and V. A. Bloomfield. 2001b. Force-induced melting of the DNA double helix 2. Effect of solution conditions. Biophys. J. 80:894-900) The force results in DNA denaturation. 10 Primers are added to the medium, and the stretching force is reduced to less than 60 pN. This allows the primers to anneal to the denatured, single-stranded DNA. DNA polymerase and nucleotides are added to the medium, and highly accurate replication is initiated by reducing the stretching force to between 0 and 30 pN. After replication is complete, a stretching force of approximately 65 pN is applied to each of the double-stranded DNA complexes, resulting in the release of single-15 stranded DNA molecules. This can be scaled up by using an array of manipulators. For example, an array such as the optical trap arrays made by Arryx, Inc. can be

While the invention has been described in detail with reference to particular embodiments thereof, it will be apparent to one skilled in the art that various changes can be made, and equivalents employed, without departing from the scope of the invention.

used.

THE CLAIMS DEFINING THE INVENTION ARE AS FOLLOWS:

- 1. A method for amplifying a nucleic acid, comprising:
 - (a) contacting one or more template strands of single-stranded nucleic acid with one or more oligonucleotide primers complementary to a portion of the one or more template strands;
 - (b) annealing at least one primer of the one or more primers to the portion of the one or more template strands to which the primer is complementary;
 - (c) contacting the one or more template strands with a nucleic acid
 polymerase and at least four different nucleoside triphosphates;
 - (d) extending the at least one annealed primer by the nucleic acid polymerase thereby forming one or more extension products bound to the one or more template strands;
 - (e) separating the one or more extension products from the one or more template strands; and
 - (f) repeating steps (a), (b), (c), (d) and (e) to amplify the nucleic acid, wherein at least one of the one or more extension products in step (e) is used as template strands in a subsequent cycle of steps (a)-(e),

wherein, for the last cycle, step (e) is optional, and wherein at least one of steps (b), or (d) comprises applying tension to the one or more template strands.

- 2. The method of claim 1, further wherein step (c) comprises applying tension to the one or more template strands.
- 3. The method of claim 1, wherein the application of tension is controlled and comprises applying tension to the one or more template strands comprises applying mechanical tension, hydrodynamic stress, or electrical field energy to the one or more template strands.

4. The method of claim 3,

wherein applying mechanical tension to the one or more template strands comprises moving a surface to which the one or more template strands are attached, or

wherein applying tension to the one or more template strands comprises applying force to one or more particles connected to one or both ends of at least one strand of the one or more template strands.

- 5. The method of claim 1, wherein applying tension to the one or more template strands comprises controlling hydrodynamic tension caused by fluid flow on the one or more template strands.
- 6. The method of claim 5, wherein the one or more template strands are anchored to a surface exposed to a fluid flow, the one or more template strands are bound to a DNA polymerase anchored to a surface exposed to a fluid flow, or the one or more template strands are retained and extended by counter-propagating fluid flows.
- 7. The method of claim 1, wherein step (b) comprises applying tension to the one or more template strands within the range selected from the ranges of about 0 to about 65 pN, about 10 to about 65 pN, and greater than 0 pN to about 45 pN.
- 8. The method of claim 1, wherein step (d) comprises applying tension to the one or more template strands and increasing the tension applied at a time corresponding to a period during which a difficult sequence on the nucleic acid is replicated.
- 9. The method of claim 8, wherein the nucleic acid comprises one or more segments of several residues which have a GC content greater than 50 percent and the nucleic acid is amplified at greater than 90 percent

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efficiency.

- The method of claim 8, wherein the nucleic acid comprises one or more 10. segments comprising a repeating unit of 8 or more base pairs and the nucleic acid is amplified at greater than 90 percent efficiency.
- 11. The method of claim 1, wherein steps (a), (b), (c), (d) and (e) are conducted at below about 65°C; or

wherein the steps (a), (b), (c), (d) and (e) are repeated at least about 50 times; or wherein a template strand of greater than about 1000 bases is amplified; or wherein the method is completed in about 1 hour or less.

- 12. A method for amplifying a nucleic acid, comprising:
 - (a) contacting one or more template strands of single-stranded nucleic acid with one or more oligonucleotide primers complementary to a portion of the one or more template strands, wherein at least one primer of the one or more primers comprises one or more complexing groups capable of binding one or more molecules bound to a substrate;
 - (b) annealing the at least one primer with the one or more complexing groups to the portion of the one or more template strands to which the primer is complementary, thereby binding the one or more template strands to the substrate;
 - (c) contacting the one or more template strands with a nucleic acid polymerase and at least four different nucleoside triphosphates;
 - (d) extending the one or more primers by the nucleic acid polymerase thereby forming one or more extension products bound to the one or more template strands;
 - (e) separating the one or more extension products from the one or more template strands; and

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- (l) repeating steps (a), (b), (c), (d) and (e) to amplify the nucleic acid, wherein, in the last cycle, step (e) is optional,
- wherein at least one of the one or more extension products in step (e) is used as template strands in a subsequent cycle of steps (a)-(e), and wherein at least one of steps (b), (d), and (e) comprises applying tension to the one or more template strands.
- 13. The method of claim 12, wherein the one or more complexing groups are activatable.
- 14. The method of claim 12, wherein the one or more complexing groups and the one or more molecules bound to the substrate are biotin and streptavidin, respectively.
- 15. The method of claim 12, wherein the one or more complexing groups comprise photoactivatable caged biotin.
- 16. A method of detecting a pathogen comprising the method of claim 1 and further comprising probing the amplified nucleic acid for the presence of a nucleotide sequence specifically associated with the pathogen.
- 17. A method for amplifying a nucleic acid, comprising:
 - denaturing a double-stranded nucleic acid molecule by applying (a) tension to one or both strands of the double-stranded nucleic acid molecule sufficient to cause separation of the double-stranded nucleic acid molecule into two single-stranded nucleic acid molecules, wherein the one or both single-stranded nucleic acid molecules to which the tension is applied becomes a template strand;
 - (b) contacting the one or both template strands with one or more oligonucleotide primers complementary to a portion of the template strands, a nucleic acid polymerase and at least four different nucleoside triphosphates;

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- (c) reducing the tension applied to the one or both template strands in step a) to permit the one or more primers to anneal to the corresponding portions of the template strands to which the one or more primers are complementary;
- (d) controlling the tension applied to the one or both template strands so as to permit extension of the one or more primers by the nucleic acid polymerase, thereby forming one or more double-stranded nucleic acid molecules in which one or more extension products are bound to the one or more template strands; and
- repeating steps (a), (b), (c), and (d) to amplify the nucleic acid,
 wherein at least one of the one or more double-stranded
 nucleic acid molecules formed in step (d) in a cycle of steps (a)-(d) is
 used as the double-stranded nucleic acid molecule in step (a) in a
 subsequent cycle of steps (a)-(d), so that at least one extension
 product is formed in step (d) is used as a template strand in a
 subsequent cycle of steps (a)-(d).
- 18. The method of claim 17, wherein at least one of applying, reducing, and controlling the tension comprises binding the nucleic acid molecules to which the tension is applied to a substrate or one or more particles or both and controlling movement of said substrate or the one or more particles or both; or

wherein at least one of applying, reducing, and controlling the tension comprises binding the nucleic acid molecules to which the tension is applied to one or more particles and controlling movement of said one or more particles by optical or magnetic manipulation.

- 19. The method of claim 17, wherein at least one of applying, reducing, and controlling the tension comprises modulating fluid flow over the nucleic acid molecules to which the tension is applied.
- 20. The method of claim 17, wherein at least one of said nucleic acid molecules

is bound by a polymerase that is bound to the substrate, and wherein fluid is flowed over the substrate.

- 21. The method of claim 17, wherein at least one of said nucleic acid molecules is bound to a substrate.
- 22. The method of claim 17, wherein at least one of said nucleic acid molecules is stretched in a fluid flow velocity gradient or by a counter propagating fluid flow.
- 23. The method of claim 17, wherein step (d) comprises increasing the tension at a time corresponding to a period during which a difficult sequence on the one or more template strands is replicated.
- 24. The method of claim 17, wherein the process is carried out at room temperature and the tension in step b) is above about 65 pN, in step c) the tension is reduced to between about 0 pN to about 60 pN, and in step d) the tension is modulated between about 0 pN to about 45 pN.
- 25. A method of performing nucleic acid amplification, comprising:
 - denaturing a double-stranded nucleic acid to produce single-stranded template strands;
 - annealing at least one primer to the template strands; (b)
 - (c) extending the at least one primer by a template-strand-driven polymerase to produce extension products, thereby forming double-stranded nucleic acid molecules; and
 - (d) repeating steps (a)-(c) to amplify the double-stranded nucleic acid,

wherein at least one of the extension products in step (c) is used as template strand in a subsequent cycle of steps (a)-(c), and

wherein step (c) comprises applying adjustable tension to the nucleic acid, said adjustable tension modulating proofreading exonuclease activity of the nucleic acid polymerase.

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26. A method for controlling the stringency of annealing of a primer to a nucleic acid strand during nucleic acid amplification, comprising:

annealing the primer to the nucleic acid strand, wherein tension is adjustably applied to said nucleic acid strand or to said primer to control the stringency of annealing of said primer to said nucleic acid strand.

27. A method for lowering the error rates of extending a primer by a nucleic acid polymerase during nucleic acid amplification, comprising:

extending the primer annealed to the nucleic acid template by a nucleic acid polymerase to produce double-stranded nucleic acid molecules, wherein tension is adjustably applied to said nucleic acid template or to said primer to lower the error rates of extending said primer by the nucleic acid polymerase.

28. A method for lowering the error rates of extending a primer on a GC-rich template by a nucleic acid polymerase during nucleic acid amplification, comprising:

extending the primer annealed to the GC-rich nucleic acid template by a nucleic acid polymerase to produce double-stranded nucleic acid molecules, wherein tension is adjustably applied to said GC-rich nucleic acid template or to said primer to lower the error rates of extending said primer on said GC-rich template.

- 29. A method of improving fidelity of nucleic acid sequencing, comprising:
 - (a) providing four samples, each sample comprising:
 - a single-stranded or a double-stranded DNA target sequence;
 - a DNA polymerase;
 - a primer complementary to the target sequence;
 - at least four different nucleoside triphosphates; and
 - a respective dideoxy nucleoside triphosphate selected from
 - among ddATP, ddCTP, ddGTP, and ddTTP;

- (b) denaturing the double-stranded DNA target sequence into singlestranded DNA target sequence;
- (c) hybridizing the primer to its complementary single-stranded DNA target sequence;
- (d) extending the primers to form double-stranded DNA extension products while adjustably applying tension to the template strand to induce proofreading exonuclease activity, thereby improving fidelity or primer extension;
- (e) repeating steps (b) through (d) to form further extension products, wherein at least one of the extension products formed in step (d) is used as a template strand in a subsequent cycle of steps (b)-(d), and;
- (f) determining the sequence of the DNA target sequence, wherein more than one of the steps (b) through (d) comprises applying tension to one or more strands of DNA target sequence.
- 30. A product produced by the method of claim 29.
- 31. A method of detecting presence of a pathogen in a biological sample, comprising:
 - (a) performing a nucleic acid amplification:
 - (i) denaturing a double-stranded nucleic acid to produce single-stranded template strands;
 - (ii) annealing at least one primer to the template strands;
 - (iii) extending the at least one primer by a template-strand-driven polymerase to produce extension products, thereby forming double-stranded nucleic acid molecules; and
 - (iv) repeating steps (i)-(iii) to amplify the double-stranded nucleic acid,

wherein at least one of steps (i)-(iii) comprises applying tension to the nucleic acid, and

(b) probing the amplified nucleic acid for the presence of a nucleotide sequence specifically associated with the pathogen,

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wherein pathogen or the nucleotide sequence are detected at concentrations lower than 1000 organisms or polynucleotides/ml.

- 32. The method of claim 31, wherein the tension applied is selected from mechanical tension, hydrodynamic tension, or electromagnetic tension, or a combination thereof.
- 33. The method of claim 31, wherein pathogen or polynucleotides are detected at concentrations lower than 100 organisms or polynucleotides/m1; at concentrations lower than 50 organisms or polynucleotides/ml; concentrations lower than 25 organisms or polynucleotides/ml; or concentrations lower than 5 organisms or polynucleotides/ml; or at a concentration of 1 organism or polynucleotide/ml.
- 34. A product produced by the method of claim 31.
- 35. A method of detecting presence of at least one specific nucleic acid sequence in a sample, comprising:
 - (a) performing a nucleic acid amplification:
 - (i) denaturing a double-stranded nucleic acid to produce singlestranded template strands;
 - (ii) annealing at least one primer to the template strands;
 - (iii) extending the at least one primer by a template-strand-driven polymerase to produce extension products, thereby forming double-stranded nucleic acid molecules; and
 - (iv) repeating steps (i)-(iii) to amplify the double-stranded nucleic acid, thereby producing amplified nucleic acid,

wherein at least one of steps (i)-(iii) comprises applying tension to the nucleic acid, and

- (b) contacting the amplified nucleic acid with at least one probe; and
- (c) detecting whether the specific nucleic sequence is in the sample by observing the presence or absence of the probe bound to the

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amplified nucleic acid.

- 36. A method of distinguishing between at least two specific nucleic acid sequences in a sample, comprising:
 - (a) performing a nucleic acid amplification:
 - (i) denaturing a double-stranded nucleic acid to produce single-stranded template strands;
 - (ii) annealing at least one primer to the template strands;
 - (iii) extending the at least one primer by a template-strand-driven polymerase to produce extension products, thereby forming double-stranded nucleic acid molecules; and
 - (iv) repeating steps (i)-(iii) to amplify the double-stranded nucleic acid, thereby producing amplified nucleic acid,

wherein at least one of steps (i)-(iii) comprises applying tension to the nucleic acid, and

- (b) contacting the amplified nucleic acid with a plurality of probes; and
- (c) detecting which of a plurality of probes is bound to the amplified nucleic acid, thereby distinguishing between at least two specific nucleic acid sequences.
- 37. A method of amplification or detection of sequences encoded on RNA, comprising:
 - (a) isolating RNA from a sample;
 - (b) performing reverse transcription thereby obtaining a corresponding cDNA template;
 - (c) amplifying the corresponding cDNA template by a method, including:
 - (i) optionally denaturing a double-stranded eDNA template to produce at least one single-stranded eDNA template strand;
 - (ii) annealing at least one primer to the at least one cDNA template strand;
 - (iii) extending the at least one primer by a template-strand-driven

polymerase to produce extension products, thereby forming double-stranded nucleic acid molecules; and

(iv) repeating steps (i)-(iii) to amplify the double-stranded nucleic acid, thereby amplifying or detecting sequences encoded on RNA,

wherein at least one of steps (i)-(iii) comprises applying tension to the nucleic acid.

38. A method for controlling denaturing of a double stranded nucleic acid during nucleic acid amplification, comprising:

adjustably applying hydrodynamic tension to said nucleic acid strand to control the denaturing of said double stranded nucleic acid.

- 39. A product of nucleic acids produced by the method of Claim 1, wherein
 - a. the average nucleic acid length can be up to 20,000 base pairs; or
 - the nucleic acid product contains less than 10⁻⁷ errors /base for normal sequences; or
 - c. the nucleic acid product contains less than 10⁻³ errors/base for difficult sequences; or
 - d. the nucleic acid product is produced by more than 100 cycles of amplification.
- 40. A method for amplifying a nucleic acid, comprising:
 - (a) contacting one or more template strands of single-stranded nucleic add with one or more oligonucleotide primers complementary to a portion of the one or more template strands;
 - (b) annealing at least one primer of the one or more primers to the portion of the one or more template strands to which the primer is complementary;
 - (e) contacting the one or more template strands with a nucleic acid polymerase and at least four different nucleoside triphosphates;
 - (d) extending the at least one annealed primer by the nucleic acid polymerase,

thereby forming one or more extension products bound to the one or more template strands;

- separating the one or more extension products from the one or more template strands; and
- (f) repeating steps (a), (b), (c), (d) and (e) to amplify the nucleic acid, wherein at least one of the one or more extension products in step (e) is used as template strands in a subsequent cycle of steps (a)-(c),

wherein, for the last cycle, step (e) is optional, and wherein tension is applied during each cycle of steps (a)-(e), and, optionally, wherein each step (a) through (e) is isothermal.

- 41. A method for improving fidelity of a nucleic acid amplification, comprising:
 - (a) contacting one or more template strands of single-stranded
 nucleic acid with one or more oligonucleotide primers
 complementary to a portion of the one or more template strands;
 - (b) annealing at least one primer of the one or more primers to the portion of the one or more template strands to which the primer is complementary;
 - (c) contacting the one or more template strands with a nucleic acid polymerase and at least four different nucleoside triphosphates;
 - (d) extending the at least one annealed primer by the nucleic acid polymerase, thereby forming one or more extension products bound to the one or more template strands;
 - separating the one or more extension products from the one or more template strands; and
 - (f) repeating steps (a) through (e) to amplify the nucleic acid,
 wherein at least one of the one or more extension products
 in step (e) is used as template strands in a subsequent cycle of

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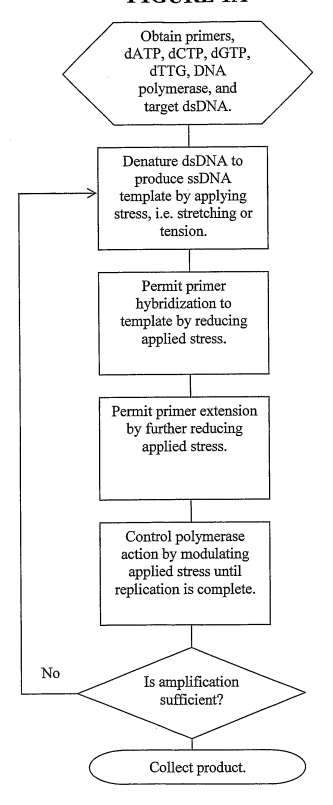
steps (a)-(e) and wherein, for the last cycle, step (e) is optional; and wherein either

tension is applied during each cycle of steps (a)-(e); or wherein tension induces proofreading exonuclease activity; or wherein tension is applied to control at least one of the step (a) through (c),

thereby improving fidelity of amplification.

- 42. The product produced by the method of Claim 41.
- 43. A method of any one of claims 1, 12, 16, 17, 25, 26, 27, 28, 29, 31, 35, 36, 37, 38, 40 or 41, substantially as herein described with reference to any example thereof.
- 44. A product of any one of claims 30, 34, 39 or 42, substantially as herein described with reference to any example thereof.

FIGURE 1A



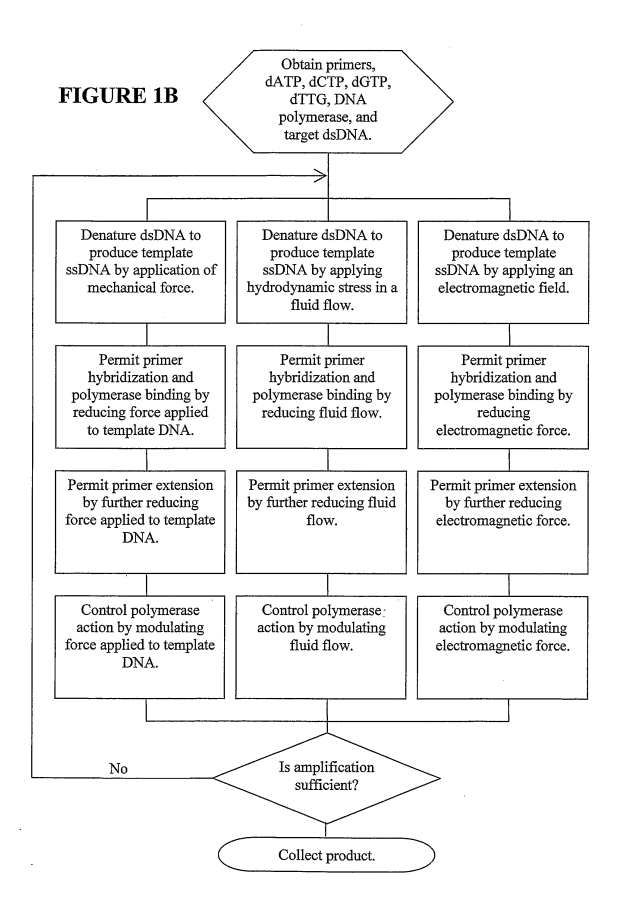


FIGURE 2A

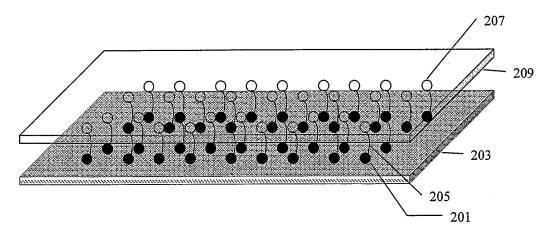


FIGURE 2B

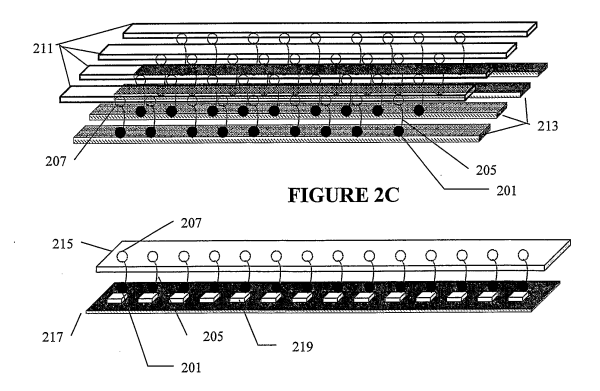
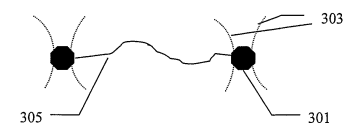
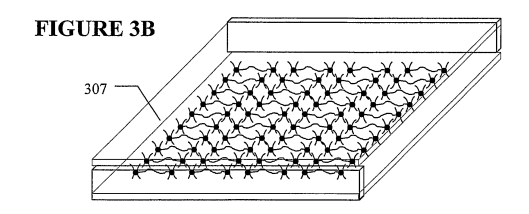
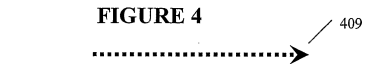


FIGURE 3A







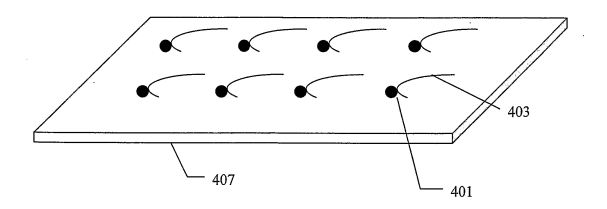


FIGURE 5A

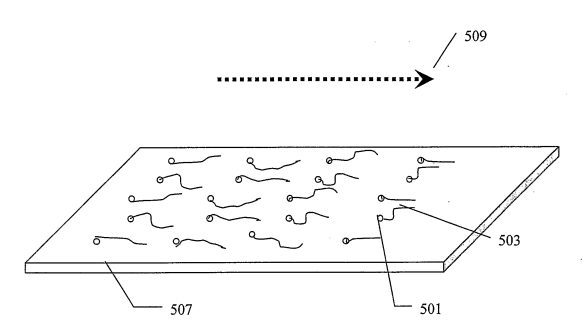


FIGURE 5B

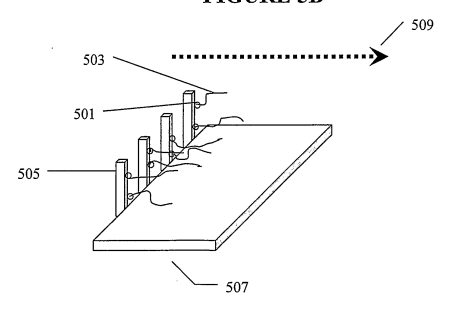
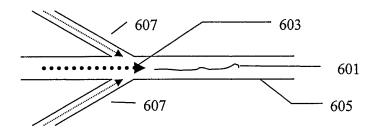
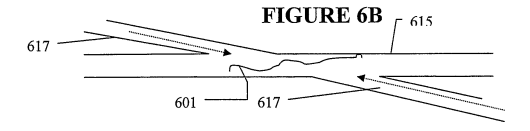


FIGURE 6A





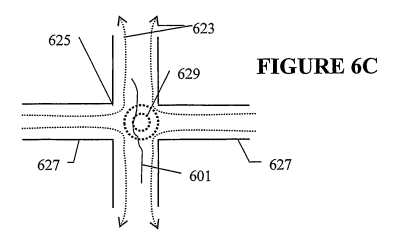


FIGURE 7A

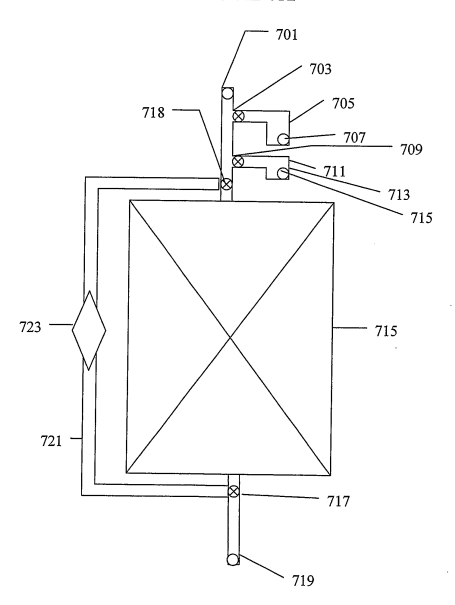


FIGURE 7B

